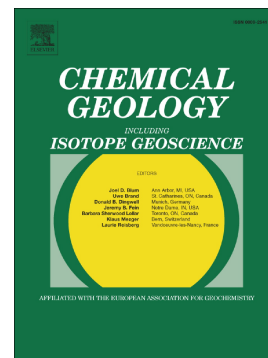


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# **The use of operationally-defined sequential Fe extraction methods for mineralogical applications: a cautionary tale from Mössbauer spectroscopy**

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## Abstract

Reactive iron minerals are crucial components of global nutrient cycles, directly controlling carbon transport and storage in marine sediments. Sequential selective extraction is frequently used for quantitatively characterising, and chemically isolating, individual Fe mineral phases. Reagent-specific mineral solubility is fundamental to the success of any sequential extraction, but is strongly affected by the varying physical and chemical morphology intrinsic to natural mineral samples. Natural sediment, rock, and soil samples often contain a mineral mixture, which further modifies solvent efficacy.  $^{57}\text{Fe}$  Mössbauer spectroscopy only probes the hyperfine interactions between next-nearest neighbouring atomic nuclei in the crystal lattice and is less affected by variation in mineral grain size and crystallinity than conventional, X-ray-based methods. In this study, we used Mössbauer spectroscopy in a novel context to cross-calibrate and optimise a popular, but frequently misused, sequential Fe extraction protocol. Our results showed that incomplete and premature removal of the target Fe minerals could occur at nearly every stage of the extraction and, in many cases, the leachate Fe content did not represent the target phase at all. Crystalline, natural siderite and amorphous, synthetic goethite were detected in the Mössbauer spectrum of the ammonium oxalate extraction for magnetite, after which all reactive Fe minerals should have been removed. Consistent with previous studies, and unlike many other clay minerals, nontronite was extracted as part of the highly reactive Fe pool, and in fact our data indicate that this mineral was extracted by the initial Na-acetate extraction that targets 'carbonate-bound Fe'. Matrix effects appeared to cause variable yield efficiencies: synthetic goethite was successfully removed when present as an individual mineral yet persisted beyond its target extraction when present in an Fe mineral mixture. Although suitable for the quantification of operationally-defined Fe pools, we caution the unverified use of sequential Fe extraction protocols for mineral specific applications. The application of sequential Fe

extractions to define the reactive Fe pools as a paleoredox proxy of depositional conditions appears relatively robust. The premature removal of 2-line ferrihydrite observed in this study (due to the use of the more aggressive Na-acetate extraction for crystalline siderite), does not limit the quantitative use of the sequential Fe extraction in ancient sediments, where such ‘easily reducible’ oxides are unlikely to persist. In contrast, attributing the outcomes of operationally-defined Fe pools to specific Fe minerals is precarious and potentially entirely erroneous. Where Fe mineral specificity or separation is required, we recommend post-extraction validation by another secondary technique. Mössbauer spectroscopy offers such a method that can independently verify extraction stages and assess mineral specificity.

**Keywords:** colloid; nanoparticle; grain size; semi-quantitative characterisation.

### HIGHLIGHTS

- The operational nature of sequential chemical extractions is frequently misunderstood when used to identify specific mineral phases, and extraction results are often unverified.
- The incomplete and premature dissolution of target minerals can occur throughout the Poulton and Canfield (2005) extraction for Fe.
- Grain size and matrix effects largely control the accuracy and the precision of many extraction stages.
- The use of Mössbauer spectroscopy (MBS) alongside sequential extractions can provide a means of quantitative Fe mineral identification.
- MBS can characterise amorphous colloidal and nanoparticulate Fe minerals.

## 1. INTRODUCTION

An estimated  $21.5 \pm 8.6$  % of organic carbon (OC) sequestered within marine sediments is bound directly to reducible Fe(III) oxides (Lalonde et al., 2012). Typically present as amorphous colloids ( $< 1 \mu\text{m}$  diameter) or nanoparticles ( $< 0.1 \mu\text{m}$  diameter), the Fe(III) oxides have a high surface area-to-volume ratio. Herein, the term “amorphous” refers to both nanoparticulate and true, non-crystalline phases. The stable Fe:OC complexes that protect OC from degradation in oxic marine sediments over million-year timescales (Estes et al., 2019; Lalonde et al., 2012) are currently attributed to the presence of fine-grained material with a large surface area for adsorption (Keil et al., 1994; Mayer, 1994; Berner, 1970) and, increasingly, the specific mineralogy of the Fe (III) oxides (Barber et al., 2017; Ransom et al., 1998). Mixtures of colloidal and nanoparticulate Fe minerals cannot be characterised by conventional analytical techniques (e.g. XRD, transmission electron microscopy), since the particles are generally too small and lack a sufficiently crystalline structure. Sequential selective leaching protocols that relate chemically-mobilised Fe to specific mineral fractions (Poulton and Canfield, 2005; Haese et al., 1997; Heron et al., 1994; Kostka and Luther, 1994; Raiswell et al., 1994; Wallmann et al., 1993; Afonso and Stumm, 1992; Canfield, 1989), are therefore used to better constrain the Fe mineralogy in modern and ancient sediments.

Sequential extraction procedures rely on the selective dissolution of a specific metal-bearing phase by particular reagents. The reagents become increasingly aggressive and less specific as the sequence progresses and the metal content of a sample is separated into “operationally-defined” groups (e.g. acetate-extractable Fe) (Kersten et al., 1997; Gobeil et al., 1995; Ure, 1991). However, sequential extraction results are often interpreted as the metal content that is bound to a particular mineral phase (e.g. carbonate-bound Fe), despite the unreliability and non-specificity reported from certain extraction stages (Eiche et al., 2010; Reinhard et al., 2009; Sulkowski and Hirner, 2006; Tlustos et al., 2005; Baeyens et al., 2003; Parat et al.,

2003; Sahuquillo et al., 1999). Such interpretations ignore the operational nature of the selective extractions and cannot assume the presence of any particular Fe mineral phase in the leachate. There has been a long-standing concern with the use of unverified extraction results to identify specific minerals, or mineral groups, in modern soils and sediments (Sanchez-Espana et al., 2019; Adamo et al., 2018; Schröder et al., 2016; Hass and Fine, 2010; Sutherland, 2010; Zimmerman and Weindorf, 2010; Bacon and Davidson, 2008; Ryan et al., 2008; Gómez-Ariza et al., 1999; Whalley and Grant, 1994; Kheboian and Bauer, 1987). However, a second, independent analytical method that is capable of characterising amorphous Fe colloids and nanoparticles is not always available.

The Poulton and Canfield (2005) sequential extraction procedure for Fe is an accessible technique that uses readily-available equipment and reagents. Since it is also capable of quantitatively processing large sample batches, it is used widely throughout the geochemical discipline. The extraction has been applied to ancient sediments in combination with several other independent proxies, to reconstruct palaeoceanic redox conditions (Hammarlund et al., 2019; Oonk et al., 2018; Raiswell et al., 2011; Reinhard et al., 2009) and constrain the timing of major biological radiations in Earth's evolutionary history (Chen et al., 2015; Dahl et al., 2010; Canfield et al., 2007). In modern sediments, the Poulton and Canfield (2005) method has been used to quantify benthic Fe fluxes in continental shelf margin sediments (Scholz et al., 2014; Homoky et al., 2013; Severmann et al., 2010) and to evaluate the importance of bioavailable Fe released from subglacial sediment sources into the open ocean (Henkel et al., 2018; Markussen et al., 2016; Raiswell et al., 2008). The Fe extraction has helped to establish the fundamental role of Fe in global S (Blonder et al., 2017; Wehrmann et al., 2017), P (Ghaisas et al., 2019; Lenstra et al., 2018; Ding et al., 2016), and C (Ma et al., 2018; Peter and Sobek, 2018; Barber et al., 2017) cycling and sequestration. Further to this, the extraction has been used to define the mechanisms for key nutrient (Li et al., 2018; Kraal et al., 2017;

Ma et al., 2017; Dijkstra et al., 2016) and toxic metal (Lynch et al., 2018; Nemati et al., 2009; Weber et al., 2009) mobilisation in freshwater sediment systems.

In their original study, Poulton and Canfield (2005) demonstrated the accurate dissolution of carbonate-associated Fe ( $\text{Fe}_{\text{CARB}}$ ; including siderite and ankerite), easily reducible Fe oxides ( $\text{Fe}_{\text{OX1}}$ ; including ferrihydrite and lepidocrocite), reducible oxides ( $\text{Fe}_{\text{OX2}}$ ; including goethite,

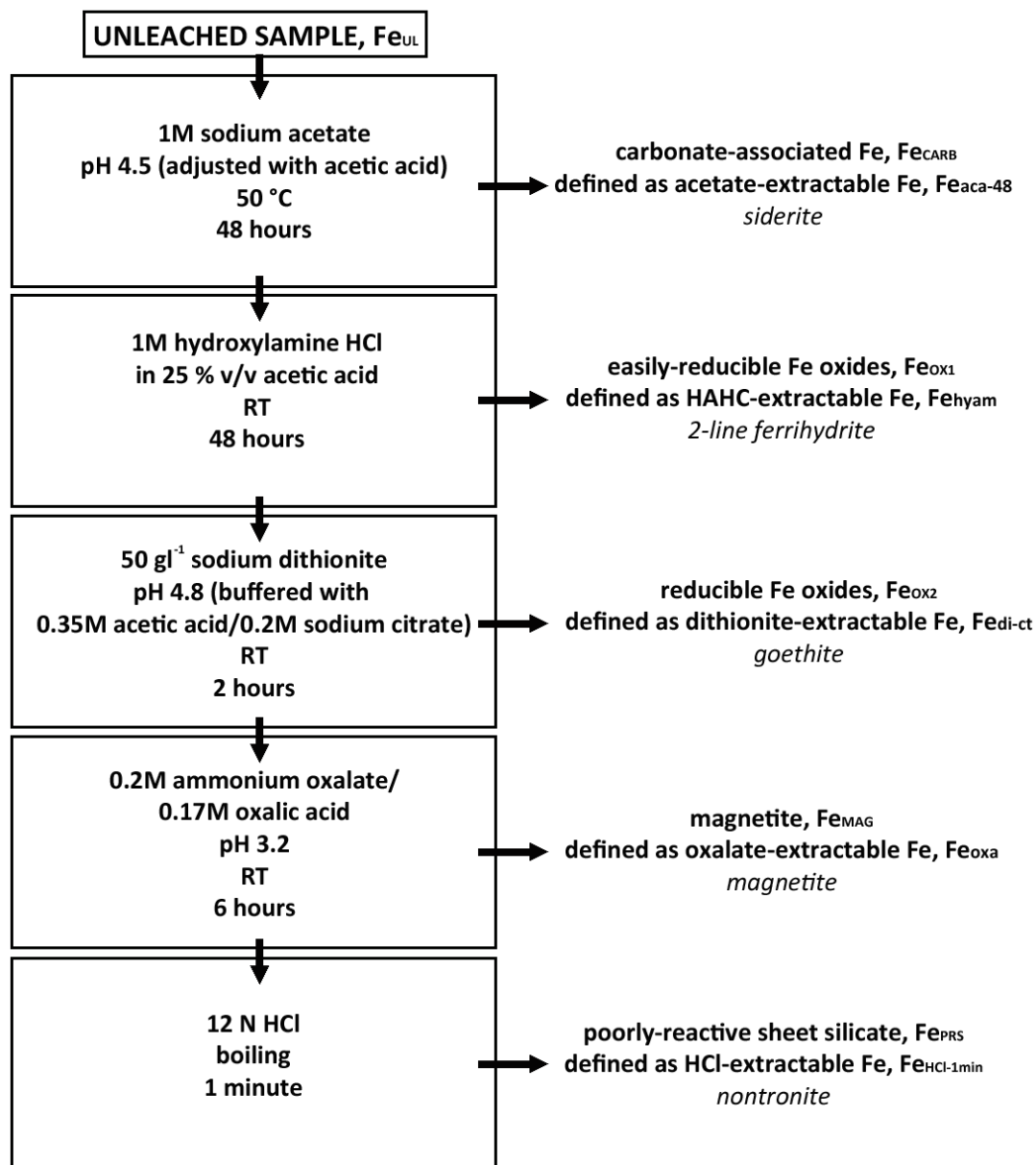


Figure 1: (1.5-column fitting image in black & white) The Poulton and Canfield (2005) sequential extraction procedure for Fe as used in this study. Each extractant is defined to reflect the operational nature of the method, after Henkel et al. (2016, 2018). The mineral selected from each of the “operationally-derived iron pools” defined by Poulton and Canfield (2005), and targeted at each stage in this study, is italicised beneath each labelled extractant. The final  $\text{Fe}_{\text{HCl-1min}}$  does not ‘target’ nontronite, but rather extracts Fe that is poorly reactive towards dissolved sulfide. RT = room temperature.

haematite, and akaganéite), magnetite, ( $\text{Fe}_{\text{MAG}}$ ), poorly-reactive sheet silicate ( $\text{Fe}_{\text{PRS}}$ ), pyrite ( $\text{Fe}_{\text{PY}}$ ), and unreactive silicate Fe ( $\text{Fe}_{\text{US}}$ ) from pure mineral phases. In many subsequent studies, the Poulton and Canfield (2005) sequential extraction for Fe has been applied as a redox indicator, to compare the sum of highly reactive Fe ( $\text{Fe}_{\text{CARB}} + \text{Fe}_{\text{OXI}} + \text{Fe}_{\text{MAG}} + \text{Fe}_{\text{PY}}$ ) with that of total Fe; in such cases, mineral specific information is not required. However, where sequential extractions are used to discern precise sample mineralogy, it is important that the extraction results are verified.

We have combined a modification of the standard Poulton and Canfield (2005) sequential extraction technique (shown in Fig. 1 and explained in detail in Section 2.4) with room temperature  $^{57}\text{Fe}$  Mössbauer spectroscopy (MBS), to semi-quantitatively verify the specific Fe minerals that are targeted at each extraction stage. MBS probes the hyperfine interactions at the  $^{57}\text{Fe}$  nucleus between the electromagnetic field of the nucleus, the electromagnetic field of the electron shell, and the electromagnetic field of the crystal lattice. Essentially, MBS utilises a measurement of a mineral directly, rather than the response of a mineral to a chemical reagent and offers a completely independent analytical method. MBS is ideally suited to the identification of reactive Fe minerals in marine sediments because 1) it can individually quantify and characterise different carbonates, iron oxides and oxyhydroxides, sulfides, sulfates, and silicates, and distinguish between the magnetic minerals magnetite and greigite (although the Mössbauer parameter space for certain minerals will overlap); 2) it requires no long range ordering of the crystal lattice, so is capable of analysing crystalline and amorphous colloids, nanoparticles (low temperature measurements are necessary to identify superparamagnetic phases in small particles), and larger phases; 3) it is only sensitive to Fe, any matrix effects caused by more abundant but less reactive mineral phases, e.g. quartz, are blended out; 4) it further determines iron oxidation states, quantifies the distribution of iron between mineral phases and oxidations states, and provides information



on the magnetic properties, spin states, coordination, bonding properties, crystallinity, and particle size.

MBS has been applied previously to verify individual extraction stages in the sequential Poulton and Canfield (2005) procedure. Following on from experiments simulating Precambrian banded iron formation diagenesis (Posth et al., 2013), the Poulton and Canfield (2005) sequential extraction procedure was used with the intent to extract individual mineral phases for subsequent isotope analysis. Isotope analysis was not carried out because MBS showed that the minerals were not separated as intended. Ferrihydrite was used as a synthetic phase at the start of the experiment discussed by Schröder et al. (2016); lepidocrocite (which is also targeted by the  $\text{Fe}_{\text{hyam}}$  stage) was not present as a starting material and did not form during the high-pressure and high-temperature experiment. After high-pressure and high-temperature treatment, MBS showed unequivocally that no ferrihydrite was present (indicating that all of it had transformed into new minerals), where the sequential extraction suggested the presence of hydroxylamine HCl-extractable Fe oxides (Schröder et al., 2016). The extracted Fe must have come from another, non-target mineral phase. MBS also revealed the presence of siderite beyond its target extraction stage, and the premature removal of magnetite (Schröder et al., 2016).

In this study we applied the Poulton and Canfield (2005) procedure, as shown in Fig. 1, to an Fe mineral mixture sediment analogue. We compared the leachate Fe content from each extraction stage with MBS analysis of the residual, leached sediment. We used a suite of natural and laboratory-synthesised Fe minerals to identify the extraction stages in which the target minerals were prematurely or incompletely dissolved. For many of our analyses, we replicated the same synthetic materials that were used to establish the original Poulton and Canfield (2005) method. The use of MBS in this study is not intended as a replacement for sequential selective extraction, rather as a means of testing the extraction method and

assessing its suitability for use in different geochemical applications. We have also adopted the extraction stage terminology defined by Henkel et al. (2018, 2016) rather than the original Poulton and Canfield (2005) definitions (Fig. 1), to exemplify the operational nature of the leaching protocol and to avoid misinterpretation of our leach results.

## 2. MATERIALS AND METHODS

### 2.1 Fe mineral preparation

One target mineral was selected from each of the “operationally-derived iron pools” defined by Poulton and Canfield (2005), excluding exchangeable metals, pyrite and unreactive silicate Fe: Ivigtut (igneous) siderite (for  $\text{Fe}_{\text{CARB}}/\text{Fe}_{\text{aca-48}}$ ), synthetic 2-line ferrihydrite (for  $\text{Fe}_{\text{OX1}}/\text{Fe}_{\text{hyam}}$ ), natural and synthetic goethite (for  $\text{Fe}_{\text{OX2}}/\text{Fe}_{\text{di-ct}}$ ), synthetic magnetite (for  $\text{Fe}_{\text{MAG}}/\text{Fe}_{\text{oxa}}$ ), and two natural nontronite specimens (for  $\text{Fe}_{\text{PRS}}/\text{Fe}_{\text{HCl-1min}}$ ) respectively represent the sodium acetate-, hydroxylamine-HCl- (HAHC-), sodium dithionite-, ammonium oxalate-, and HCl-extractable fractions (Fig. 1). It is important to note that the final  $\text{Fe}_{\text{PRS}}$  stage is not mineral specific. Unlike the preceding stages which are designed to target specific operationally-defined mineral groups (Poulton and Canfield, 2005),  $\text{Fe}_{\text{PRS}}$  removes Fe that is poorly reactive towards dissolved sulphide (Raiswell and Canfield, 1996). Raiswell et al. (1994) demonstrated the quantitative extraction of Fe from both oxide/oxyhydroxide and silicate phases using the boiling HCl method of Berner (1970). The use of  $\text{Fe}_{\text{HCl-1min}}$  in this study, is simply to test how effectively it removes the nontronite and any residual Fe mineral phases left behind by the earlier stages. The target minerals were chosen based on their importance in modern sediment studies, ease of MBS identification, and availability. Pyrite is easy to identify using X-ray based techniques and was excluded from this study because it does not face the same issues of characterisation as the more amorphous Fe oxides

and oxyhydroxides. Unreactive silicate was also excluded because the bioreactive Fe minerals are the focus of this study.

A crystalline sample of siderite from Ivigtut, Greenland, and a powdered sample of natural goethite (unknown locality) were obtained from the teaching collection, School of Geosciences, University of Edinburgh (UK). The natural goethite is hereafter referred to as UoE goethite. The powdered UoE goethite was confirmed as such by reflected light microscopy (before powdering). One natural nontronite (Hungary) sample was kindly donated by the National Museums Scotland Collection Centre (Edinburgh, UK); this sample is henceforth referred to as NMS nontronite. A sample of NAu-2 nontronite, originally collected from Uley Mine, South Australia, was purchased from The Clay Minerals Society (Virginia, U.S.A.). Freshly exposed surfaces of all, natural minerals were selected beneath a hand lens, then ground prior to analysis; the maximum and average grain sizes of each natural mineral are provided in Table 1.

Synthetic samples of 2-line ferrihydrite, goethite, and magnetite were prepared from the standard Cornell and Schwertmann (2003) methods, identical to those used to establish the original Poulton and Canfield (2005) method. To synthesise the 2-line ferrihydrite, 500 ml 0.1 M iron(III) nitrate was stirred continuously during the addition of 330 ml 1 M potassium hydroxide, until the pH reached 7 – 8. After centrifugation, the suspension was rapidly transferred to sealed sections of dialysis tubing and submerged in ultra-pure H<sub>2</sub>O (18.2 MΩ.cm) baths at room temperature, to remove electrolytes. 8.5 g of 2-line ferrihydrite was produced by freeze-drying the final, dark red-brown product. Ferrihydrite precipitation was also the first stage involved during goethite synthesis: 180 ml 5 M potassium hydroxide was added to 100 ml 1 M iron(III) nitrate solution. The suspension was diluted to 2 L with ultra-pure H<sub>2</sub>O (18.2 MΩ.cm) and sealed in a polypropylene bottle in a preheated, 70 °C oven for 60 hours. The final yellow-coloured goethite was then washed repeatedly using vacuum

filtration and finally dried at 50 °C, to produce approximately 8 g of goethite. For the magnetite synthesis, a 240 ml 3.33 M potassium hydroxide/0.27 M potassium nitrate solution was added dropwise to 560 ml of a 0.3 M iron(II) sulfate solution preheated to 90 °C. All solutions involved in the magnetite synthesis were sparged with N<sub>2</sub> for 2 hours before use. The suspension was heated for a further 60 minutes under constant agitation before the black, magnetite precipitate (approximately 5 g) could be washed, filtered, and dried. All synthetic minerals were dried then ground prior to analysis; see Table 1 for the synthetic Fe mineral maximum and average grain and minimum crystallite sizes. The characterisation of all Fe minerals, natural and synthetic, was performed by both XRD (Appendix A) and MBS (Appendix B).

## 2.2 XRD

The mineralogy of the natural Ivigtut siderite and synthetic magnetite samples was analysed using a PANalytical X'Pert PRO diffractometer at the National Museums Collection Centre (Edinburgh, UK) using CuK $\alpha$  radiation, and identified by comparison with the International Centre for Diffraction Data database, PDF-2, using X'pert High Score software. The mineralogy of the synthetic 2-line ferrihydrite, synthetic goethite, and natural NMS and NAu-2 nontronite samples was analysed likewise at Cardiff University with a monochromator to reduce Fe fluorescence.

## 2.3 Mineral grain size

At least three photomicrographs were made of each ground Fe mineral under transmitted light prior to analysis. Examples of these can be found in Fig. 2. The photomicrographs were individually processed using Fiji (Schindelin et al., 2012). In Fiji, the images were individually binarized and threshold adjusted (manually, using the sliding scale in the threshold tool) to define every visible mineral grain. Grain clusters were erased from the original photomicrograph where no clear border between separate mineral grains could be

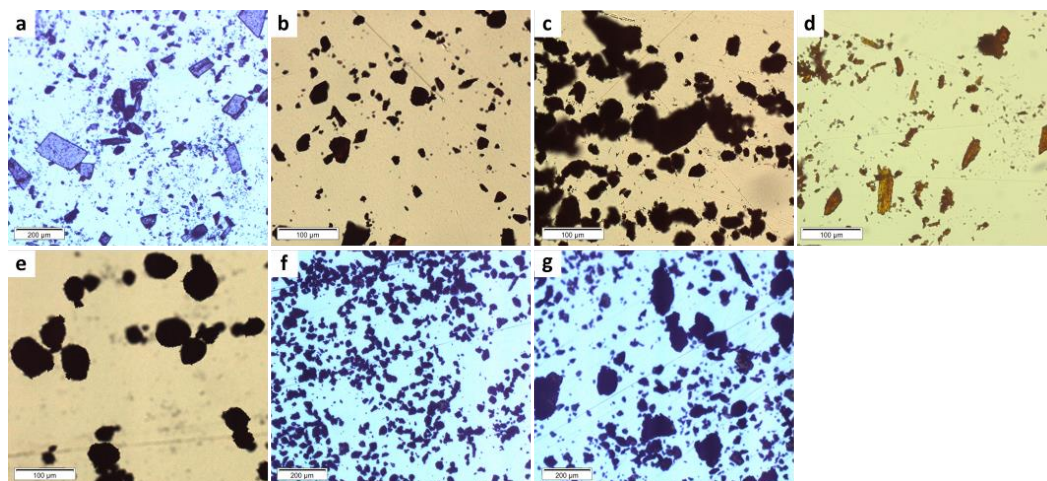


Figure 2: (1.5-column fitting image, in full colour online only) Photomicrographs of the Fe minerals used in this study under transmitted light. a) Ivigtut siderite (natural), b) 2-line ferrihydrite (synthetic), c) goethite (synthetic), d) UoE goethite (natural), e) magnetite (natural), f) NMS nontronite (natural), and g) NAu-2 nontronite.

seen. All sizes (from 0 to infinity) and shapes (with circularity set from 0 – 1) of particles were analysed, exclusive of particles overlapping the edge of the photomicrograph, which were discarded. The maximum Feret particle dimensions are reported in Table 1. Grains less than 5  $\mu\text{m}$  in diameter could not be reliably defined; these data were also discarded.

XRD data for each Fe mineral was combined with the Scherrer equation (Equation 1), as described in Guilbaud et al. (2010), on each identified  $2\theta$  peak, to roughly estimate the average crystallite (sub-micrometre particle) diameter of the smallest (and most reactive) particles that were excluded from the Fiji image analysis. The Scherrer equation defines the lower bound of crystallite size to the broadening of a specific  $2\theta$  peak in an XRD spectrum:

Equation 1

$$L = K\lambda(\beta\cos\theta)^{-1}$$

where  $L$  is crystallite diameter, the Scherrer constant ( $K$ ) used is 0.91 assuming spherical particles (Brindley, 1980),  $\lambda$  is the wavelength of the X-ray (0.154 nm for  $\text{CuK}\alpha$ ),  $\beta$  is the full width of the peak at half maximum (FWHM, in radian) and  $\theta$  (in radian) is the angle of the peak.

		Ivigtut siderite (natural)	2-line ferrihydrite (synthetic)	goethite (synthetic)	UoE goethite (natural)	magnetite (synthetic)	NMS nontronite (natural)	NAu-2 nontronite (natural)
Particle diameter ( $\mu\text{m}$ )	Maximum	$177 \pm 18$	$236 \pm 12$	$161 \pm 16$	$172 \pm 17$	$157 \pm 16$	$119 \pm 12$	$208 \pm 21$
	Average	$16 \pm 2$	$19 \pm 1$	$28 \pm 3$	$26 \pm 3$	$25 \pm 3$	$25 \pm 3$	$28 \pm 3$
	$n_p$	3425	885	835	4024	3346	4484	3780
Crystallite diameter (nm)	Minimum	53	0.6	14	15	12	3	2
	Average	57	0.8	21	25	17	9	6
	$n_c$	2	2	4	4	4	4	5

Table 1: Maximum and average Feret diameter ( $\mu\text{m}$ ), and number of sampled grains ( $n_p$ ) used to measure Fe mineral grain size. All mineral grain size analyses were performed using the Fiji Image J software. Minimum and average crystallite diameters (nm), and number of 2 $\theta$  peaks ( $n_c$ ) used to estimate crystallite size. All crystallite diameters were estimated from the Scherrer equation (Equation 1); a precision of  $\pm 9$  nm was determined from 5 repeat measurements of the same sample.

Table 1 reports the average and maximum grain size values calculated using Fiji, and the estimated minimum and average crystallite diameters using the XRD data combined with the Scherrer equation (Equation 1). The reported errors for the Fiji-analysed grain sizes represent the minimum size at which particles could be identified for the specific magnification: for photomicrographs magnified by 2X – 4X and 10X – 20X, this is  $\pm 10\%$  and  $\pm 5\%$  respectively. The precision on  $L$  was  $\pm 9$  nm, as calculated from 5 repeat measurements of the UoE goethite sample. Grain size distribution is also compared between the two goethite and between the two nontronite samples used in this study, in Fig. 3.

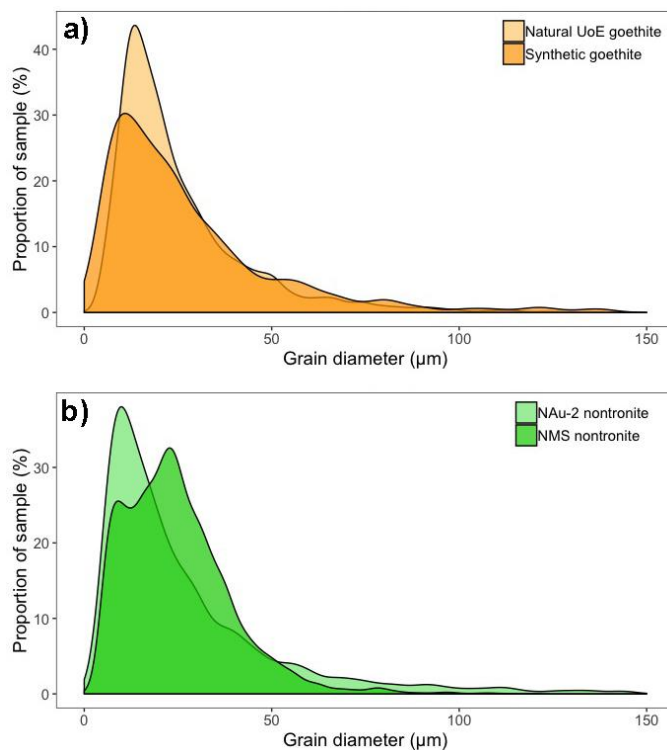


Figure 3: (single-column fitting image, in full colour online only) Distribution of mineral grain size throughout unleached, ground, and homogenised mineral samples. a) natural UoE and synthetic goethite, and b) NMS nontronite and NAu-2 nontronite. Mineral grain size analyses were performed using the Fiji Image J software.

## 2.4 Sequential Fe extraction

A mixed Fe mineral standard (CARB-2: approximately 30 % siderite, 20 % ferrihydrite, 20 % goethite, 15 % magnetite, and 15 % nontronite) was made to represent an example of the Fe mineral assemblage (not the entire mineral assemblage) that may occur in natural marine

sediments, using just one mineral from each target Fe mineral phase in the extraction sequence. Each of the selected Fe minerals and the mixed Fe mineral standards were run through the full sequential extraction protocol shown in Fig. 1. Each stage of the same protocol was also performed non-sequentially on the individual Fe mineral targeted by that particular stage: whereby only  $\text{Fe}_{\text{aca-48}}$  was run for siderite; only  $\text{Fe}_{\text{hyam}}$  was run for 2-line ferrihydrite; only  $\text{Fe}_{\text{di-ct}}$  was run for goethite; and  $\text{Fe}_{\text{oxa}}$  was run for magnetite. The  $\text{Fe}_{\text{HCl-1min}}$  stage is not designed to specifically target nontronite.  $\text{Fe}_{\text{HCl-1min}}$  was therefore run individually for nontronite to test the effectiveness of nontronite dissolution by a 1-minute boil in HCl (as described below). Each non-sequential, single mineral extraction was performed in triplicate; the average value from each extraction is presented in Table 2.

Total Fe was extracted from approximately 0.1 g of the ground, dried, and unleached individual and mixed Fe mineral samples using a three-stage digestion procedure: 4 ml aqua regia; 2 ml HF; 2 ml  $\text{HNO}_3$ . The digestions were each performed in Teflon beakers left on a hotplate at 90 °C for 24 hours, dried completely, and cooled before adding the next reagent. All samples were finally redissolved in 2 %  $\text{HNO}_3$  for dilution and Fe concentration analysis by atomic absorption spectrometry (AAS).

The modified Poulton and Canfield (2005) Fe extraction scheme was run as described in Fig.

1. All citrate-buffered dithionite solutions were prepared using fresh dithionite powder immediately before each batch of  $\text{Fe}_{\text{di-ct}}$  extractions. The ' $\text{Fe}_{\text{aca-48}}$ ' and ' $\text{Fe}_{\text{HCl-1min}}$ ' terminology denotes variations to the standard ' $\text{Fe}_{\text{aca}}$ ' and ' $\text{Fe}_{\text{HCl}}$ ' Poulton and Canfield (2005) methods, which we employed during the first and last stages of our sequential protocol (Fig. 1). We used the longer (48, rather than 24 hours) and hotter (50 °C, rather than room temperature) Na-acetate leach to target our highly-crystalline Ivigtut siderite sample, as recommended in Poulton and Canfield (2005). However, this more aggressive leach is rarely used in sequence on modern sediments since it is likely to cause premature leaching of



subsequent Fe minerals, such as ferrihydrite, as is observed in this study. The  $\text{Fe}_{\text{aca-48}}$  extraction is more readily applied to ancient sediments, in which ferrihydrite is rarely found (Cornell and Schwertmann, 2003), and premature removal of this easily-reducible Fe oxide is not an issue. All extraction stages were performed in sealed centrifuge tubes on a shaker table under oxic conditions, except  $\text{Fe}_{\text{HCl-1min}}$ . In the final  $\text{Fe}_{\text{HCl-1min}}$  extraction, 12 N HCl was heated on a hotplate in a glass beaker and, once boiling, the sample was added and stirred into the reagent. After boiling for exactly 1 minute, the solution was quenched immediately with ultra-pure  $\text{H}_2\text{O}$  (18.2 M $\Omega$ .cm), then decanted for centrifugation. At the end of each extraction stage (Fig. 1) the suspension was centrifuged at 6000 rpm for 10 minutes, which allowed the leachate to be carefully pipetted from the residual solid sample. The leachate was subsampled and diluted in 2 %  $\text{HNO}_3$  ready for Fe analysis by AAS. 0.05 – 0.15 g of washed (three times with ultra-pure  $\text{H}_2\text{O}$  (18.2 M $\Omega$ .cm) and re-centrifuged) and dried (at 30 °C overnight) sample was removed for analysis by MBS.

A sample mass : reagent volume ratio (S:R) of <2.5 (Thompson et al., 2019) was applied to each stage of the extraction to avoid saturation of the reagent by the sample. The S:Rs used to extract the Fe mineral phases in this study were much lower than that recommended by Poulton and Canfield (2005) for natural sediment: 0.1 – 0.2 g : 10 ml. Since our experiment required the removal of 40 – 60 mg after each stage for MBS analysis, all extractions were performed in triplicate to ensure sufficient residual material was available for each subsequent MBS subsampling, while maintaining an optimal S:R ratio. Following each extraction, the material was combined and homogenised for MBS analysis, then divided back into triplicate to maintain a suitably-low S:R. As residual material was removed for MBS analysis, the S:R was gradually reduced throughout the sequence: ~2.5 in  $\text{Fe}_{\text{aca-48}}$ , ~2.1 in  $\text{Fe}_{\text{hyam}}$ , ~1.7 in  $\text{Fe}_{\text{di-ct}}$ , ~1.3 in  $\text{Fe}_{\text{oxa}}$ , and ~0.8 in  $\text{Fe}_{\text{HCl-1min}}$ . Thompson et al. (2019) recommend a lower S:R of ~0.75 for  $\text{Fe}_{\text{di-ct}}$ , than was used herein; the possible implications

of this are discussed in Section 3.1. No residue was left for MBS analysis following the stages that completely extracted the target mineral. The total digestions were run separately, rather than as the final stage in the extraction sequence. The entire procedure (Fig. 1) was repeated 3 times for each individual mineral and the mixed Fe mineral standard (each performed in triplicate) to initially produce a total of 9 leachate results, and three batches of residual material per sample; the number of samples ( $n$ ) from each extraction stage continually reduced throughout the sequence as more material was removed for MBS. The average values and  $n$  from each extraction stage run in sequence for the individual Fe minerals and the mixed Fe mineral standards (CARB-2 and CARB-3) are respectively presented in Tables 3 and 4.

## 2.5 Atomic Absorption Spectroscopy (AAS)

The Fe concentration in the final leachate and digest solutions was determined on a Unicam 989 AA Spectrometer at the University of Stirling (UK) and calibrated using matrix-matched single element Fe standards. The diluted solutions were run in a random sequence and all measurements were blank and drift corrected. Measured concentrations of the certified reference material (CRM) SdAR-H1 (blended metalliferous sediment) were accurate to  $\pm 5\%$  of the value reported by the International Association of Geoanalysts ([http://iageo.com/wp-content/uploads/2017/11/SdAR-H1\\_RM\\_data\\_sheet-1.pdf](http://iageo.com/wp-content/uploads/2017/11/SdAR-H1_RM_data_sheet-1.pdf)). Analytical precision was  $< 6\%$  RSD, from replicate analysis of randomly chosen samples and the SdAR-H1 CRM. The Fe concentrations extracted by single stage individual mineral extractions only, the full sequence (see Fig. 1) of individual mineral extractions, and mixed Fe mineral sequential extractions is respectively reported in Tables 2, 3 and 4.

## 2.6 Mössbauer Spectroscopy (MBS)

The presence, absence, and relative proportion of specific Fe mineral phases in leached sample residues was confirmed by room temperature  $^{57}\text{Fe}$  MBS (Gütlich and Schröder,

2012), using a standard transmission Mössbauer spectrometer (Wissel, Germany) with a  $^{57}\text{Co}$  in Rh matrix radiation source in constant acceleration mode, at the University of Stirling (UK). 0.05 – 0.15 g of dried, homogenised sample was loaded into sealed acrylic discs with a circular cross section of approximately  $1\text{ cm}^2$ . Troughs in transmission spectra represent absorption maxima; the area beneath each subspectrum represents the relative contribution of the specific Fe mineral phase to the observed/fitted MBS absorption spectrum (Figs. 4 – 5).

The MBS spectra were analysed using the Voigt-based fitting routine (Rancourt and Ping, 1991) as implemented in the Recoil software (University of Ottawa, Canada). Following the Voigt-based fitting, we compared hyperfine parameters (including isomer shift ( $\delta$ ) in  $\text{mm s}^{-1}$ , quadrupole splitting ( $\Delta\text{EQ}$ ) in  $\text{mm s}^{-1}$ , and internal magnetic field ( $B_{\text{hf}}$ ) in T) of each subspectrum with those of library reference spectra. All MBS data can be found in tables C1 – C4 of the Appendix.

### 3. RESULTS

#### 3.1 Single and sequential extraction of individual Fe mineral samples

$\text{Fe}_{\text{hyam}}$  and  $\text{Fe}_{\text{oxa}}$  were the only stages to fully extract the target Fe mineral phase from the pure mineral samples. The extraction behaviour of the single Fe minerals differed between different natural samples, and between natural and synthetic variants of the same mineral.

The igneous Ivigtut siderite was particularly resistant to extraction by sodium acetate (Table 2). Less than 15 % of the total siderite Fe was collected by the individual  $\text{Fe}_{\text{aca-48}}$  stage (Table 2), even when a 48-hour leach at pH 4.5 and 50 °C was applied, as recommended by Poulton and Canfield (2005) for crystalline siderite samples. MBS also identified siderite in the residual material of each extraction stage up to and including  $\text{Fe}_{\text{oxa}}$  (Table 3) before it was fully removed in  $\text{Fe}_{\text{HCl-1min}}$ ; although 23 – 44 % of the siderite was dissolved as  $\text{Fe}_{\text{hyam}}$  (Table 3).

Sample	Target extraction	Total Fe available for extraction	Fe extracted in target stage	Post-extraction MBS
Ivigtut siderite (natural)	Fe <sub>aca-48</sub>	46.9 ± 2.1 (n = 3)	3.9 ± 0.2 (n = 3)	P
2-line ferrihydrite (synthetic)	Fe <sub>hyam</sub>	47.2 ± 2.1 (n = 3)	41.6 ± 1.8 (n = 2)	A
goethite (synthetic)	Fe <sub>di-ct</sub>	63.2 ± 2.8 (n = 3)	60.1 ± 2.6 (n = 3)	A
UoE goethite (natural)	Fe <sub>di-ct</sub>	56.6 ± 2.5 (n = 3)	23.8 ± 1.0 (n = 3)	P
magnetite (synthetic)	Fe <sub>oxa</sub>	72.2 ± 3.2 (n = 3)	75.5 ± 3.3 (n = 3)	A
NMS nontronite (natural)	Fe <sub>HCl-1min</sub>	24.5 ± 1.1 (n = 3)	12.5 ± 0.5 (n = 3)	P
NAu-2 nontronite (natural)	Fe <sub>HCl-1min</sub>	19.3 ± 0.8 (n = 3)	1.8 ± 0.1 (n = 3)	P

Table 2: Average Fe contents (wt.%) for each individual extraction stage compared with the average total Fe (wt.%) dissolved in the bulk HF / HNO<sub>3</sub> digest. Each extraction stage was performed non-sequentially using the single target Fe mineral for each stage: only siderite was run through Fe<sub>aca-48</sub>; only 2-line ferrihydrite was run through Fe<sub>hyam</sub>; only goethite was run through Fe<sub>di-ct</sub>; only magnetite was run through Fe<sub>oxa</sub>; and only nontronite was run through Fe<sub>HCl-1min</sub>. MBS analyses were conducted on residual material left from the target extractions; the presence (P) or absence (A) of each individual mineral in the post-extraction residue is shown in the final column. 'n' = number of samples.

Sample	Target extraction	Total Fe available for extraction	Fe extracted in Fe <sub>aca-48</sub>	Post-Fe <sub>aca-48</sub> MBS	Fe extracted in Fe <sub>hyam</sub>	Post-Fe <sub>hyam</sub> MBS	Fe extracted in Fe <sub>di-ct</sub>	Post-Fe <sub>di-ct</sub> MBS
Ivigtut siderite (natural)	Fe <sub>aca-48</sub>	46.9 ± 2.1 (n = 3)	1.5 ± 0.1 (n = 9)	P +	15.3 ± 0.7 (n = 8)	P +	2.6 ± 0.1 (n = 6)	P -
2-line ferrihydrite (synthetic)	Fe <sub>hyam</sub>	47.2 ± 2.1 (n = 3)	6.8 ± 0.3 (n = 7)	P	44.2 ± 1.9 (n = 6)	A	ND	A
goethite (synthetic)	Fe <sub>di-ct</sub>	63.2 ± 2.8 (n = 3)	BDL (n = 9)	P	± 0.0 (n = 9)	P -	54.6 ± 2.4 (n = 7)	A
UoE goethite(natural)	Fe <sub>di-ct</sub>	56.6 ± 2.5 (n = 3)	0.0 ± 0.0 (n = 9)	P	0.1 ± 0.0 (n = 8)	P	35.8 ± 1.6 (n = 4)	P
magnetite (synthetic)	Fe <sub>oxa</sub>	72.2 ± 3.2 (n = 3)	0.0 ± 0.0 (n = 9)	P	0.6 ± 0.0 (n = 8)	P	31.4 ± 1.4 (n = 5)	P
NMS nontronite (natural)	Fe <sub>HCl-1min</sub>	24.5 ± 1.1 (n = 3)	± 0.0 (n = 9)	P -	0.9 ± 0.0 (n = 6)	P -	13.8 ± 0.6 (n = 4)	P
NAu-2 nontronite (natural)	Fe <sub>HCl-1min</sub>	19.3 ± 0.8 (n = 3)	± 0.0 (n = 9)	P	1.0 ± 0.0 (n = 6)	P	9.6 ± 0.4 (n = 6)	P

Table 3 Average Fe contents (wt.%) of sequential mineral leachates, compared with the average total Fe (wt.%) dissolved in the bulk HF/HNO<sub>3</sub> digest. All extraction stages (Fig. 1) were run sequentially on each individual Fe mineral. Fe<sub>HCl-24</sub> (redefined from “unreactive silicate” or “Fe<sub>US</sub>”, (Poulton and Canfield, 2005)) refers to the unreactive Fe pool that remains unextracted after the sample is ashed at 450 °C for 8 hours and near-boiled in 6 N HCl for 24 hours. The average values shown in Tables 2 – 4 represent data ranges that are often highly variable, due to the operational nature and consequent unreliable reproducibility of sequential extraction protocols. MBS analyses were conducted on residual material left from the target extractions; the presence (P) or absence (A) of each individual mineral in the post-extraction residue is shown in the final column; ‘+’ and ‘-’ indicate a noticeable increase and decrease, respectively, in the relative amount of a mineral compared with the previous extraction stage. ‘P’ without ‘+’ or ‘-’ indicates no noticeable change. The cumulative total Fe adds together all of the sequentially-extracted Fe, for comparison with the total Fe initially available for extraction in the unleached Fe mineral phases (errors calculated by adding the individual errors in quadrature). BDL = below detection limit, ND = no data.

Sample	Target extraction	Total Fe available for extraction	Fe extracted in Fe <sub>oxa</sub>	Post-Fe <sub>oxa</sub> MBS	Fe extracted in Fe <sub>HCl-1min</sub>	Post- Fe <sub>HCl-1min</sub> MBS	Fe extracted in Fe <sub>HCl-24</sub>	Post-Fe <sub>HCl-24</sub> MBS	Cumulative total Fe
Ivigtut siderite (natural)	Fe <sub>aca-48</sub>	46.9 ± 2.1 (n = 3)	2.4 ± 0.1 (n = 2)	P +	32.7 ± 1.4 (n = 2)	A	ND	A	54.5 ± 1.6
2-line ferrihydrite (synthetic)	Fe <sub>hyam</sub>	47.2 ± 2.1 (n = 3)	ND	A	ND	A	ND	A	51 ± 1.9
goethite (synthetic)	Fe <sub>di-ct</sub>	63.2 ± 2.8 (n = 3)	ND	A	ND	A	ND	A	54.6 ± 2.4
UoE goethite (natural)	Fe <sub>di-ct</sub>	56.6 ± 2.5 (n = 3)	0.6 ± 0.0 (n = 2)	P	27.0 ± 1.6 (n = 2)	P -	ND	A	63.5 ± 2.3
magnetite (synthetic)	Fe <sub>oxa</sub>	72.2 ± 3.2 (n = 3)	44.8 ± 2.0 (n = 1)	A	ND	A	ND	A	76.8 ± 2.6
NMS nontronite (natural)	Fe <sub>HCl-1min</sub>	24.5 ± 1.1 (n = 3)	1.5 ± 0.1 (n = 3)	P	1.6 ± 0.0 (n = 1)	P	0.9 ± 0.0 (n = 1)	A	18.8 ± 1.9
NAu-2 nontronite (natural)	Fe <sub>HCl-1min</sub>	19.3 ± 0.8 (n = 3)	0.7 ± 0.0 (n = 4)	P -	2.9 ± 0.1 (n = 3)	P -	5.1 ± 0.2 (n = 1)	A	19.3 ± 3.1

Table 3 continued.

	Extraction stage	Total Fe available for extraction	Fe extracted in target stage	Post-extraction MBS				
				Ivigtut siderite	2-line ferrihydrite	goethite	magnetite	nontronite
CARB-2	Fe <sub>UL</sub>	49.0 ± 2.1 (n = 3)		P	P	P	P	P
	Fe <sub>aca-48</sub>	15.1 ± 0.6	14.8 ± 0.6 (n = 9)	<u>P</u> -	P -	P	P	P - (S)
	Fe <sub>hyam</sub>	10.7 ± 0.4	3.9 ± 0.4 (n = 8)	P	<u>A</u>	P +	P +	P + (S)
	Fe <sub>di-ct</sub>	10.6 ± 0.4	25.5 ± 1.1 (n = 6)	P +	A	<u>P</u> -	P	P (S)
	Fe <sub>oxa</sub>	6.3 ± 0.3	24.0 ± 1.1 (n = 4)	P	A	P	<u>A</u>	A
	Fe <sub>HCl-1min</sub>	6.3 ± 0.3	38.4 ± 1.7 (n = 1)	A	A	A	A	<u>A</u>
CARB-3	Fe <sub>UL</sub>	46.8 ± 2.1 (n = 3)		P	P	P	P	P
	Fe <sub>aca-48</sub>	14.5 ± 0.6	14.9 ± 0.7 (n = 9)	<u>P</u>	P -	P	P	P -
	Fe <sub>hyam</sub>	9.8 ± 0.4	3.9 ± 0.2 (n = 7)	P	<u>A</u>	P	P	P
	Fe <sub>di-ct</sub>	10.3 ± 0.4	30.0 ± 1.3 (n = 6)	P +	A	<u>P</u> -	P -	P +
	Fe <sub>oxa</sub>	6.1 ± 0.3	18.8 ± 0.8 (n = 5)	P	A	P -	<u>A</u>	P
	Fe <sub>HCl-1min</sub>	6.1 ± 0.3	31.5 ± 1.4 (n = 3)	P -	A	A	A	<u>P + (S)</u>
	Fe <sub>HCl-24</sub>		4.9 ± 0.2 (n = 1)	A	A	A	A	A

Table 4: Average Fe contents (wt.%) of sequential mixed mineral leachates, compared with the average total Fe (wt.%) dissolved in the bulk HF/HNO<sub>3</sub> digest. The total Fe available for each extraction stage was calculated by dividing the total Fe<sub>UL</sub> content by the known proportion of each mineral mixed into the CARB-2 and CARB-3 samples. All extraction stages (Fig. 1) were run sequentially on the CARB-2 and CARB-3 Fe mineral mixtures. Fe<sub>HCl-24</sub> (redefined from “unreactive silicate” or “Fe<sub>US</sub>”, (Poulton and Canfield, 2005)) refers to the unreactive Fe pool that remains unextracted after the sample is ashed at 450 °C for 8 hours and near-boiled in 6 N HCl for 24 hours. No ‘unreactive silicate’ Fe minerals, as defined by Poulton and Canfield (2005),

were used in this study (hence there is no Fe available for the  $\text{Fe}_{\text{HCl-24}}$  extraction stage), however the  $\text{Fe}_{\text{HCl-24}}$  stage was required to dissolve the residual nontronite following  $\text{Fe}_{\text{HCl-1min}}$ . The average values reported here often represent a wide data range (low minimum and high maximum values) that exemplify the effects of non-target mineral extraction as discussed in the text. MBS analyses were conducted on residual material left from the target extractions. The presence (P) or absence (A) of each individual mineral in the post-extraction residue is reported alongside any noticeable changes in the shape of an Fe mineral peak and a consequent modification of the mineral-specific parameters (S); '+' and '-' indicate a noticeable increase and decrease, respectively, in the relative mineral proportion compared with the previous extraction stage. 'P' without '+' or '-' indicates no noticeable change. The target mineral (underlined> of each extraction stage is highlighted in bold where the target extraction is successful, and italicised where incomplete. BDL = below detection limit, ND = no data.



Synthetic 2-line ferrihydrite was completely removed during its target ( $\text{Fe}_{\text{hyam}}$ ) stage (Table 2 and Table 3). However, 15 % of the ferrihydrite Fe was leached in  $\text{Fe}_{\text{aca-48}}$  (Table 3), likely due to the use of the extended Na-acetate leach in sequence (as discussed in Section 4.1).

More than 90 % of the total Fe available in the synthetic goethite was removed by  $\text{Fe}_{\text{di-ct}}$  when this stage was run individually (Table 2), and more than 80 % was removed when  $\text{Fe}_{\text{di-ct}}$  was run sequentially (Table 3). According to the Mössbauer spectra that accompany the sequential extraction of synthetic goethite, the abundance of synthetic goethite Fe was not affected by  $\text{Fe}_{\text{aca-48}}$  and was slightly lowered for  $\text{Fe}_{\text{hyam}}$ . Synthetic goethite was removed completely by  $\text{Fe}_{\text{di-ct}}$  – the target stage (Table 3). There was insufficient residual material for MBS analysis, following both the individual and sequential extraction of synthetic goethite (Tables 2 and 3, respectively).  $\text{Fe}_{\text{di-ct}}$  was therefore considered effective when synthetic goethite was the only mineral present.

Natural UoE goethite was more resistant to the target extraction than its synthetic form (Table 2). Only 36 – 47 % of the total Fe in the UoE goethite was recovered during the individual  $\text{Fe}_{\text{di-ct}}$  stage and the UoE goethite remained clearly visible in the residual material MB spectrum (Table 2). Up to 82 % of the total Fe was, however, recovered by the sequential  $\text{Fe}_{\text{di-ct}}$  stage of the natural UoE goethite extraction (Table 3); all remaining UoE goethite Fe was recovered during  $\text{Fe}_{\text{HCl-1min}}$ . MBS identified UoE goethite in the residual material of each sequential extraction stage, including  $\text{Fe}_{\text{HCl-1min}}$  (Table 3), although the abundance of natural goethite was noticeably reduced by  $\text{Fe}_{\text{HCl-1min}}$ .

AAS leachate and MBS residue results suggest that both the individual (Table 2) and sequential (Table 3)  $\text{Fe}_{\text{oxa}}$  extractions effectively targeted all of the synthetic magnetite Fe, although up to 66 % was solubilised during the previous, sequential  $\text{Fe}_{\text{di-ct}}$  stage (Table 3).

Mössbauer spectra of the individual (Table 2) and sequential (Table 3)  $\text{Fe}_{\text{HCl-1min}}$  extractions show that successive stages of the extraction procedure dissolved an increasing amount of both natural nontronite samples, with the final  $\text{Fe}_{\text{HCl-1min}}$  stage solubilising the majority of the available nontronite Fe. The  $\text{Fe}_{\text{HCl-1min}}$  extraction is not specific to nontronite but should dissolve the remaining Fe that is poorly reactive towards sulphide (Raiswell and Canfield, 1996). The individual  $\text{Fe}_{\text{HCl-1min}}$  stage yielded 48 – 55 % of the available NMS Fe and just 9 % of the available NAu-2 Fe (Table 2). 47 – 67 % of the NMS nontronite Fe and 47 – 53 % of the NAu-2 nontronite Fe was removed by the sequential  $\text{Fe}_{\text{di-ct}}$  stage (Table 3). Nontronite removal during  $\text{Fe}_{\text{di-ct}}$  was likely responsible for the low Fe yield recovered from the residual NMS (< 6 %) and NAu-2 (< 17 %) nontronite samples during the sequential  $\text{Fe}_{\text{HCl-1min}}$  stage (Table 3).

### 3.2 Sequential extraction of mixed Fe mineral samples

The efficiency with which individual extraction stages dissolved their target minerals was further complicated by the presence of other minerals. Every stage, excluding the final  $\text{Fe}_{\text{HCl-1min}}$ , exhibited the removal of non-target Fe minerals that should not have been affected until later on in the leach sequence.  $\text{Fe}_{\text{hyam}}$  and  $\text{Fe}_{\text{oxa}}$  were, again, the only stages to completely remove their target minerals when present in an Fe mineral mixture. Following the observed incomplete extraction of natural goethite Fe, compared with the successful removal of synthetic goethite and the extraction of natural NMS nontronite Fe during the individual  $\text{Fe}_{\text{di-ct}}$  stage (Tables 2 and 3), another mixed Fe mineral standard, CARB-3, was created to further test yield efficiency. CARB-3 included Ivigtut siderite, synthetic 2-line ferrihydrite, and synthetic magnetite in the same proportions as CARB-2, but the 20 % synthetic goethite and 15 % natural NMS nontronite were respectively replaced with 20 % natural UoE goethite and 15 % natural NAu-2 nontronite.

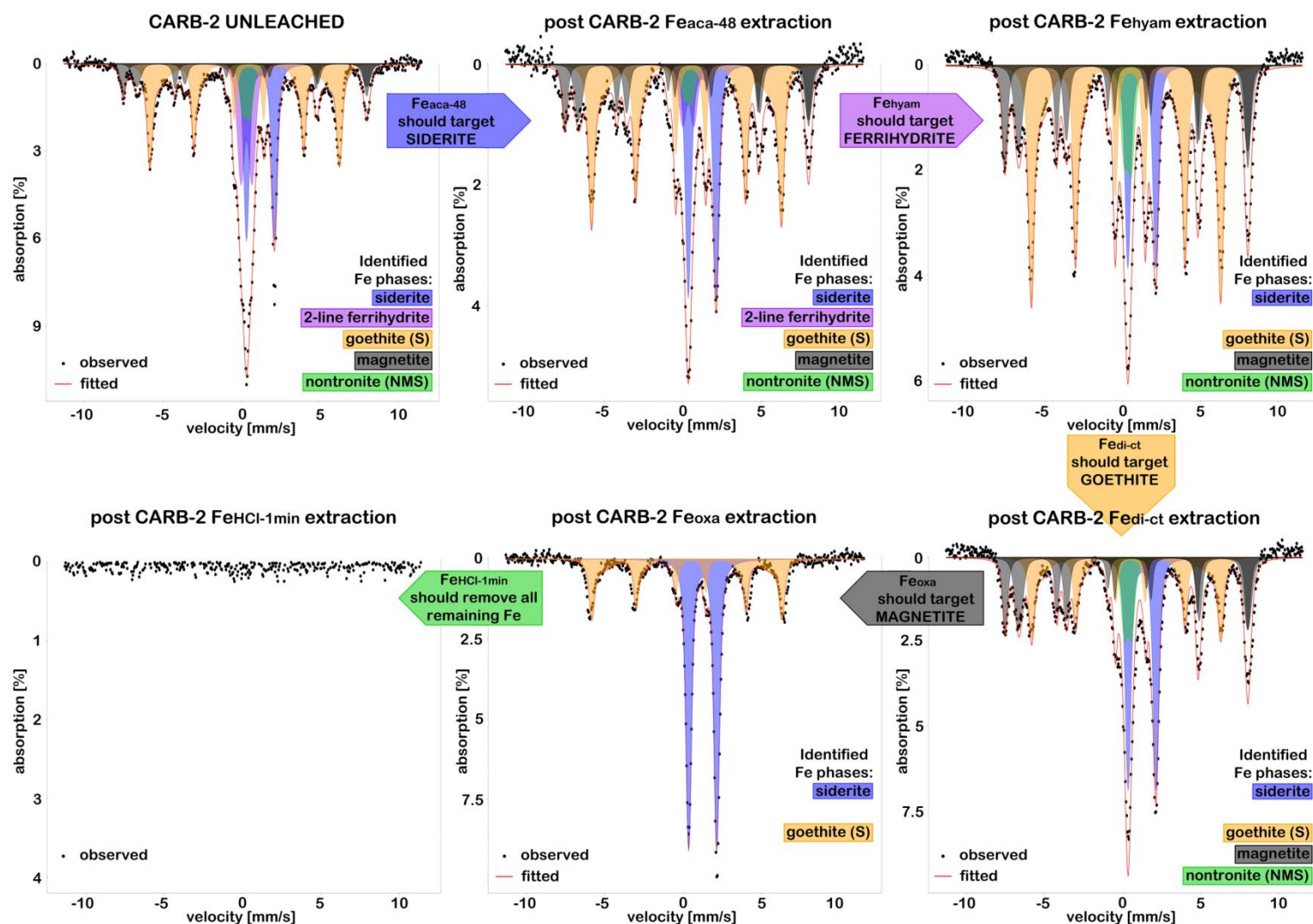


Figure 4: (2-column fitting image, in full colour online and print) Comparison of MBS spectra from the unleached and leached CARB-2 mixed mineral standard after each stage in the full sequential extraction. No Fe was present following the final  $\text{Fe}_{\text{HCl-1min}}$  stage.

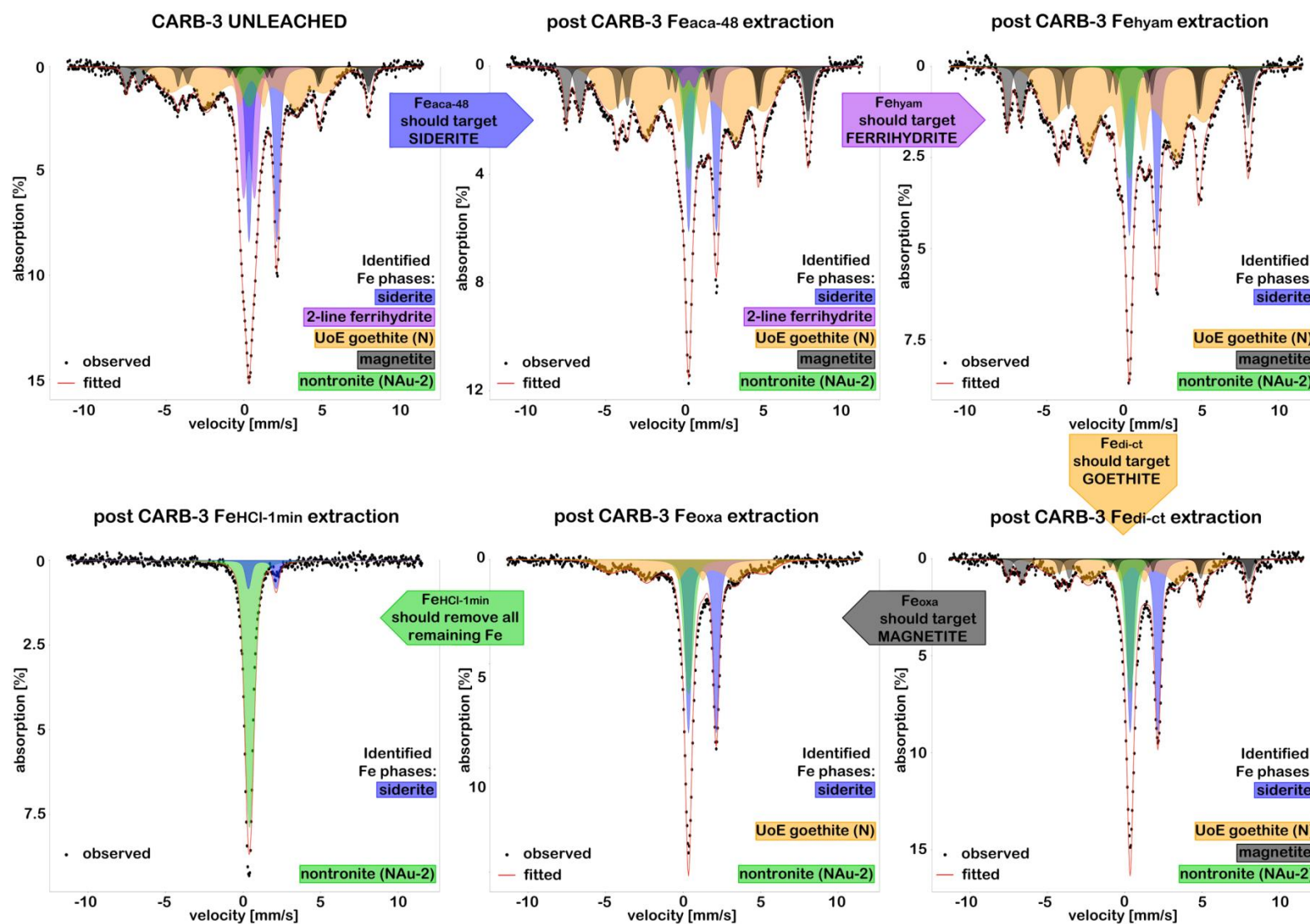


Figure 5: (2-column fitting image, in full colour online and print) Comparison of MBS spectra from the unleached and leached CARB-3 mixed mineral standard after each stage in the full sequential extraction.

The unleached CARB-2 and CARB-3 samples respectively contained 26 % and 27 % Ivigtut siderite. 26 – 35 % of the total Fe available in the unleached CARB-2 and CARB-3 samples was removed during the  $\text{Fe}_{\text{aca-48}}$  stage (Table 4), which implies that all siderite Fe was successfully removed by its target extraction. MBS analysis of the accompanying  $\text{Fe}_{\text{aca-48}}$  residue, clearly identified the presence of siderite in both CARB-2 (Fig. 4) and CARB-3 (Fig. 5) samples after the target extraction stage. In fact, siderite continued to dominate sample composition until it was completely removed during  $\text{Fe}_{\text{HCl-24}}$  (a stage that is usually only required for unreactive silicate extraction). Although siderite should be the only mineral targeted by Na acetate, the relative proportion of 2-line ferrihydrite and NMS nontronite (included in CARB-2) was noticeably reduced (Fig. 4). NAu-2 nontronite (included in CARB-3) was unaffected by  $\text{Fe}_{\text{aca-48}}$  (Fig. 5).

Synthetic 2-line ferrihydrite comprised 24 % of the unleached CARB-2 sample and 14 % of the unleached CARB-3 sample.  $\text{Fe}_{\text{hyam}}$  removed 15 – 20 % of the total Fe available in CARB-2, and 8 – 9 % of the total Fe available in CARB-3 (Table 4). MBS confirmed the complete removal of all remaining 2-line ferrihydrite Fe during the target  $\text{Fe}_{\text{hyam}}$  stage (and the removal of some 2-line ferrihydrite during  $\text{Fe}_{\text{aca-48}}$ ) in both CARB-2 (Fig. 4) and CARB-3 (Fig. 5).

Synthetic goethite accounted for 28 % of the unleached CARB-2 sample, while 34 % of CARB-3 consisted of natural UoE goethite. The CARB-2  $\text{Fe}_{\text{di-ct}}$  leachate contained a highly variable 23 – 71 % of the total available Fe (Table 4), which suggests incomplete synthetic goethite dissolution at the lower bound, and dissolution of both synthetic goethite and other mineral phases at the higher bound. The average dithionite-associated Fe content in the  $\text{Fe}_{\text{di-ct}}$  CARB-2 residue was noticeably reduced in comparison to that of the previous  $\text{Fe}_{\text{hyam}}$  stage (Fig. 4), but synthetic goethite was clearly identified in the residual material of the target  $\text{Fe}_{\text{di-ct}}$  and the following  $\text{Fe}_{\text{oxa}}$  stages (Fig. 4), which supports the incomplete dissolution of the target mineral during  $\text{Fe}_{\text{di-ct}}$ . The CARB-3  $\text{Fe}_{\text{di-ct}}$  leachate conversely contained 43 – 52 % of

the total Fe available in the unleached CARB-3 sample (Table 4), up to 18 % more Fe than was actually available from the natural UoE goethite component. From the individual mineral extraction results (Table 3) and modified parameters in the CARB-3  $\text{Fe}_{\text{hyam}}$  and  $\text{Fe}_{\text{di-ct}}$  MBS spectra (Table A1), it is likely that the overestimated  $\text{Fe}_{\text{di-ct}}$  yield (in both CARB-2 and CARB-3) came from the dissolution of magnetite Fe and NAu-2 nontronite Fe in sodium dithionite. Following the removal of the majority of the UoE goethite during  $\text{Fe}_{\text{di-ct}}$ , siderite and nontronite dominated the residual CARB-3 material (Fig. 5).

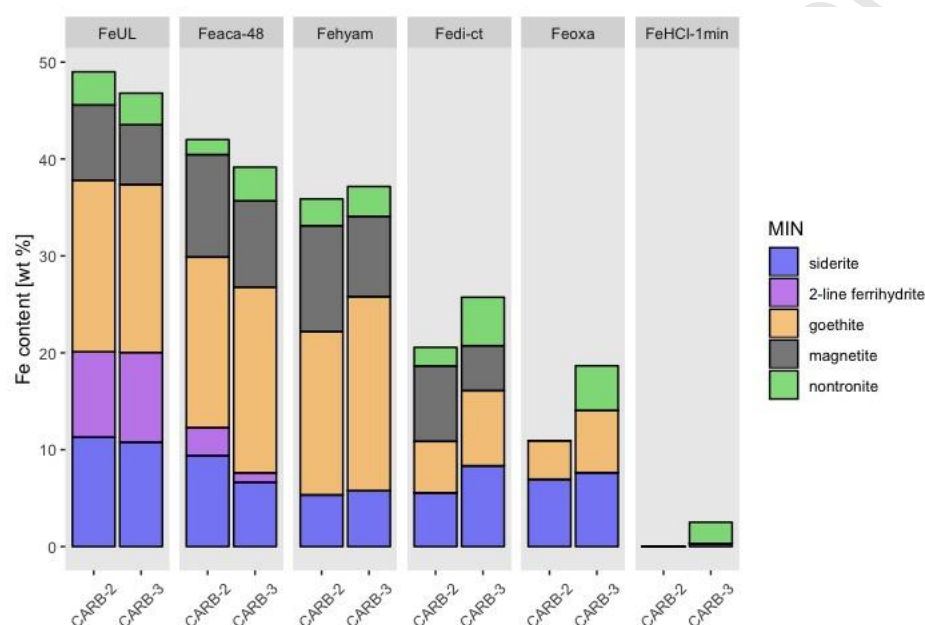


Figure 6: (1.5-column fitting image, in full colour online and print) Fe content in CARB-2 and CARB-3 throughout the extraction sequence. This figure shows a semi-quantitative estimation of the average Fe content (wt %) extracted from all minerals at each stage of the sequential leach for the mixed Fe mineral samples, CARB-2 and CARB-3. Results combined the known quantity of Fe released in the leachate at each extraction stage (measured by AAS) and the known proportion of minerals identified (by MBS) in the residual material.

All magnetite Fe present in the unleached CARB-2 and CARB-3 samples (13 – 14 %) was extracted during  $\text{Fe}_{\text{oxa}}$  (Table 4), and no magnetite was detected by MBS in the residual CARB-2 (Fig. 4) or CARB-3 (Fig. 5) material following the target extraction. Magnetite Fe was therefore effectively targeted by  $\text{Fe}_{\text{oxa}}$ , but the  $\text{Fe}_{\text{oxa}}$  leachate Fe content far exceeded that provided by the dissolution of magnetite Fe alone: up to over 60 % in both CARB-2 and CARB-3 (Table 4). The simultaneous removal of NMS nontronite Fe in CARB-2 (Fig. 4),



and natural UoE goethite Fe in CARB-3 (Fig. 5), during  $\text{Fe}_{\text{oxa}}$ , explains the presence of excess Fe in these samples. Residual siderite and synthetic goethite (from the target  $\text{Fe}_{\text{aca-48}}$  and  $\text{Fe}_{\text{di-ct}}$  stages, respectively) appeared unaffected by  $\text{Fe}_{\text{oxa}}$  in CARB-2 and remained in the residual material following the ammonium oxalate dissolution (Fig. 4); the same was true of siderite and NAu-2 nontronite in CARB-3 (Fig. 5).

All Fe minerals were removed from CARB-2 by  $\text{Fe}_{\text{HCl-1min}}$  (Fig. 4) and the residual material was bleached white (identified via XRD as quartz that was originally present in the UoE goethite and both nontronite samples). Almost all of the siderite was finally removed from CARB-3 during  $\text{Fe}_{\text{HCl-1min}}$ , although a small signal still remained in the Mössbauer spectrum (Fig. 5). The lingering Fe mineral signature post  $\text{Fe}_{\text{HCl-1min}}$  in CARB-3 (Fig. 5) was that of the NAu-2 nontronite, which appears more persistent than the NMS nontronite (possibly due to a modification of the original  $\text{Fe}_{\text{HCl}}$  Berner (1970) method). The NAu-2 nontronite was completely removed from CARB-3 by the final 24-hour boiling HCl stage, which was designed to leach the unreactive sheet silicates (Poulton and Canfield, 2005).

### 3.3 Mineral grain size and distribution

Maximum and average particle diameters, and minimum and average crystallite diameters for all the Fe minerals used in this study are shown in Table 1. Ivigtut siderite was the most crystalline of the Fe minerals used herein (Fig. 1) with the smallest average grain size.

However, the most reactive components of a sample are the crystallite grains, due to their larger surface area in comparison to the larger-diameter particles. Therefore, it is unsurprising that siderite proved particularly resistant to dissolution, since the minimum size of the siderite crystallites was more than twice that of all the other Fe minerals in this study. The natural UoE goethite had a larger maximum diameter but a smaller average diameter than the synthetic goethite and was more efficiently dissolved during its target  $\text{Fe}_{\text{di-ct}}$  stage (Table 2). NMS nontronite had both a smaller maximum and a smaller average grain size than the NAu-

2 nontronite and was much more efficiently dissolved during its target  $\text{Fe}_{\text{HCl-1min}}$  stage (Table 2). There was little difference in crystallite particle diameter between the two nontronite samples.

## 4. DISCUSSION

### 4.1 Sequential extraction limitations

Meaningful comparison between sequential extraction experiments in different (and even within the same) laboratories, is extremely difficult, due to i) the use and modification of different extraction schemes, which can produce contrasting results for identical samples (Tlustos et al., 2005); ii) the lack of either a universal standard or use of verified individual mineral samples in all studies; and iii) verification that the extracted amount of Fe comes from the intended target mineral phases. Poor siderite recovery by sequential extraction procedures is a known issue (see references in Table 5), with yields ranging from < 2 % to 100 % using a variety of methods (Table 5). The recovery of other carbonates is similarly variable and rarely reliably quantitative: 100 % for calcite; 98 % for ankerite; 45 – 95 % for dolomite; 31 % for  $\text{MnCO}_3$  and 12 % for rhodochrosite (Liu et al., 2018; Morera-Chavarria et al., 2016; Raiswell et al., 1994).

MBS analysis of post-extract substrate in this study, revealed that siderite was not only incompletely extracted during its target  $\text{Fe}_{\text{aca-48}}$  stage (as previously noted by Reinhard et al., 2009) but persisted throughout the entire procedure (Figs. 4 – 5). Even using Poulton and Canfield's (2005) extended  $\text{Fe}_{\text{aca-48}}$  procedure for crystalline siderite (48 hours at 50 °C, rather than 24 hours at room temperature), we could only remove 3 – 8 % of the total Ivigtut siderite Fe (with an average grain diameter of 16  $\mu\text{m}$ , but a maximum diameter of 177  $\mu\text{m}$ , measured by Fiji grain size analysis post pulverisation), which was significantly less than the 95 – 100 % yield of Roxbury and Biwabik siderite (both crushed to <63  $\mu\text{m}$ ) reported by



Poulton and Canfield (2005). It is possible that the reduced efficiency of  $\text{Fe}_{\text{aca-48}}$  for the Ivigtut siderite, in comparison to that of the Roxbury or Biwabik siderite, was caused by the presence of larger grains in the Ivigtut sample (Table 1 and Fig. 2a).

The  $\text{Fe}_{\text{aca-48}}$  extraction was further complicated by the apparent recovery of 30 % of the total CARB-2 and CARB-3 Fe content (Table 4), both of which contain 30 % siderite. The proportions of nontronite and 2-line ferrihydrite (Table 3) were also reduced in the CARB-2 (Fig. 4) and CARB-3 (Fig. 5)  $\text{Fe}_{\text{aca-48}}$  residue. The leachate Fe concentration implied complete siderite removal in the target  $\text{Fe}_{\text{aca-48}}$  stage by, coincidentally, dissolving the same amount of Fe from 2-line ferrihydrite and nontronite instead (Table 3, Figs. 4 – 5). As mentioned in Section 2.4, the  $\text{Fe}_{\text{aca-48}}$  leach is only normally used in sequence on ancient sediments, where ferrihydrite is unlikely to be present. The removal of 2-line ferrihydrite was therefore not an unexpected result when using the more aggressive  $\text{Fe}_{\text{aca-48}}$  leach. Although significant nontronite removal has been observed during  $\text{Fe}_{\text{di-ct}}$  (Raiswell et al. 1994), its extraction during  $\text{Fe}_{\text{aca-48}}$  was unexpected, and may again be due to the longer, 48-hour leach at 50 °C. Our main concern here was that, in isolation, the  $\text{Fe}_{\text{aca-48}}$  leach appeared to be working, when clearly MBS shows that this was not the case.

2-line ferrihydrite was successfully removed by its target  $\text{Fe}_{\text{hyam}}$  stage, when present as an individual mineral and as part of a mixture. The CARB-2 and CARB-3  $\text{Fe}_{\text{hyam}}$  leachate yielded less Fe than was expected, due to the removal of 2-line ferrihydrite during the previous,  $\text{Fe}_{\text{aca-48}}$  stage.

The apparent recovery of 25 – 30 % (Table 4) of the total Fe available in CARB-2 and CARB-3 (both of which contain approximately 20 % goethite) during  $\text{Fe}_{\text{di-ct}}$ , similarly implied that all of the goethite Fe was dissolved (along with the partial dissolution of an/other

Extraction method	Yield	Complications	Reference
	95 – 100 %	Premature ferrihydrite	
	$\text{Fe}_{\text{aca-48}}$ leachate [Fe] compared with $\text{Fe}_{\text{TOTAL}}$ [Fe] via ICP-AAS	(3 %) and lepidocrocite (1 %) removal	(Poulton and Canfield, 2005)
1 M Na acetate pH 4.5 (acetic acid) 50 °C 48 hours	80 – 85 %	Incomplete siderite removal, 5 – 10 % passed onto following extraction stage	(Oonk et al., 2017)
	3 – 8 % $\text{Fe}_{\text{aca-48}}$ leachate [Fe] compared with $\text{Fe}_{\text{TOTAL}}$ [Fe] via ICP-AAS	Incomplete siderite removal Nontronite removal	This study
1 M Na acetate pH 5.0 (acetic acid) room temperature 5 hours	< 2 %	Efficiency of carbonate removal is strongly dependent on sample:reagent ratio	(Liu et al., 2018)
cold 10 % HCl room temperature 24 hours	0 – 16.3 %	Premature removal of 'easily reducible' oxides and silicates (NQ)  Residual siderite partially removed by $\text{Fe}_{\text{oxa}}$ extraction	(Raiswell et al., 2011; Reinhard et al., 2009)
0.5 M HCl 15 hours	NQ	Minute amounts of siderite remain	(Schröder et al., 2016)
chloroacetate/maleate buffer solutions pH ~ 0 80 °C 150 minutes	100 %	Requires at least 10 mg carbonate  Not tested for use in sequential extraction (single stage only)	(Morera-Chavarria et al., 2016)

Table 5: Quantitative efficiency of modified  $\text{Fe}_{\text{aca}}$  extractions for siderite, compared with the original Poulton and Canfield (2005) method. NQ = not quantified.

non-target mineral/s). According to the accompanying MBS spectrum, goethite was clearly present in the  $\text{Fe}_{\text{di-ct}}$  residue, and that of the following  $\text{Fe}_{\text{oxa}}$  extraction (Figs. 4 – 5). It is possible that a higher S:R ratio of 1.7 used in this study during  $\text{Fe}_{\text{di-ct}}$ , compared with the < 0.75 recommended by Thompson et al. (2019), may have been partially responsible for some of the observed incomplete natural goethite extraction. However, the more successful extraction of synthetic goethite at the same S:R (Table 3), implies that other factors (e.g. grain size and matrix effects) were more likely to have caused the variable yields. Claff et al. (2010) reported the remains of > 25 % goethite in residual samples even after an 8-hour extraction and concluded that the dithionite extraction could not dissolve high concentrations of crystalline Fe oxides (including akaganéite and haematite). Non-specificity and the post-extraction re-adsorption of analytes onto residual solids can also cause variable results depending on the substrate used (Kim et al., 2015; Bacon and Davidson, 2008; Hanahan, 2004). The almost complete removal of synthetic goethite Fe during its target  $\text{Fe}_{\text{di-ct}}$  stage when run as an individual mineral (Table 2), but persistence in the residual CARB-2 mixture (Fig. 4), suggests that matrix effects may be responsible for the variable goethite yields.

Magnetite was successfully targeted by  $\text{Fe}_{\text{oxa}}$  (Figs. 4 – 5) but the AAS leachate results reveal that 28 – 69 % of the total magnetite Fe was also recovered during  $\text{Fe}_{\text{di-ct}}$  (Table 3). The amount of magnetite dissolved in sodium dithionite in this study was much higher than the 5 – 7 % reported by Poulton and Canfield (2005) but matched that of Henkel et al. (2016) at 32 – 52 %. Premature magnetite dissolution was also observed by Schröder et al. (2016). All of the Poulton and Canfield (2005) and some of the Henkel et al. (2016)  $\text{Fe}_{\text{oxa}}$  experiments, used synthetic magnetite samples that were synthesised after Cornell and Schwertmann (2003).

Arbitrary behaviour of both nontronite samples was observed throughout the entire procedure. As individual minerals, 50 – 56 % of the total NMS and N<sub>Au</sub>-2 nontronite Fe contents were recovered as dithionite-extractable Fe (Table 3); the other non-target stages

yielded negligible ( $< 6\%$ ) amounts (Table 3). Although this was higher than the  $29\%$   $\text{Fe}_{\text{di-ct}}$  recovery reported by Raiswell et al. (1994), the dissolution of nontronite Fe during  $\text{Fe}_{\text{di-ct}}$  was anticipated. However, MBS spectra of the individual mineral and the mixed Fe mineral CARB-2 samples, showed that NMS nontronite was partially removed during  $\text{Fe}_{\text{aca-48}}$  and  $\text{Fe}_{\text{hyam}}$ , before its complete dissolution in  $\text{Fe}_{\text{oxa}}$  (Fig. 4). Dissolution and structural modification of nontronite in Na acetate (Jaisi et al., 2008), HAHc (Ryan et al., 2008), sodium dithionite (Jaisi et al., 2008), and ammonium oxalate (Wu et al., 2012) is known and attributed to the reduction of structural Fe(III) (Borggaard, 1988). Conversely, the persistent presence of NAu-2 at the end of each extraction, including the  $\text{Fe}_{\text{HCl-1min}}$  stage, highlights the operational nature of sequential extractions and their critical dependence on the specific mineralogical composition of a sample. The Fe concentration measured during  $\text{Fe}_{\text{aca-48}} - \text{Fe}_{\text{oxa}}$  may therefore be overestimated by the inclusion of non-target nontronite. The extra Fe recovered from the CARB-2 and CARB-3 samples during  $\text{Fe}_{\text{oxa}}$  (Table 4) likely came from the nontronite that was removed in  $\text{Fe}_{\text{oxa}}$ .

The changing shape of the NMS nontronite spectrum within the CARB-2 sample after each consecutive extraction stage (Fig. 4), can be explained by the presence of Fe(III) in both the octahedral and tetrahedral sheets of the nontronite crystal lattice. Significant ( $> \sim 20\%$ ) nontronite dissolution can cause irreversible dehydroxylation (Fialips et al., 2002) and subsequent di-/trioctahedral substitution (Manceau et al., 2000) and partitioning of the reagent-extracted Fe(II) into surface complexation and other reactive sites (Jaisi et al., 2008). Preferential reduction of the dioctahedral sheet Fe(III) would result in a crystal structure dominated by the remaining trioctahedral sheet Fe(III), and vice versa, changing the MBS parameters and observed spectrum accordingly. Combined AAS leachate and MBS solid residue observations for CARB-2, showed that approximately  $45\%$  of the NMS nontronite Fe was removed during  $\text{Fe}_{\text{aca-48}}$  (Fig. 6), which alone would be responsible for structural

modifications to the lattice structure. Re-adsorption of the extracted Fe(II) onto the nontronite lattice may also explain why the changing shape of the NMS nontronite peak in the CARB-2 Mössbauer spectra between the Fe<sub>aca-48</sub> and Fe<sub>hyam</sub> extraction stages, was not also accompanied by increased Fe concentrations in the NMS and NAu-2 leachates.

The recovery of synthetic goethite during Fe<sub>di-ct</sub> in this study was greater than that of the natural UoE goethite (Table 2). Sequential extraction procedures are generally applied on the assumption that they target well-defined mineral phases, but natural samples often contain a complex mixture of mineral phases that are not chemically or physically distinct (Kheboian and Bauer, 1987). For example, the Fe speciation of natural mineral samples used in sequential extraction procedures is often unknown, as is the interaction between the reactant and specific solid Fe phases within the mineral matrix. However, this is exactly the information that MBS can provide, to ensure the reliable comparison of data between different sequential extraction experiments.

Poor reproducibility and non-selectivity of reactive Fe phases by wet chemical extraction has been well known for some time, leading to the development of more aggressive dissolution methods (e.g. for siderite, Table 5). Such alternative methods prove effective when used as individual leaches but compromise subsequent extraction stages when used in sequence (Raiswell et al., 1994). One such example is the room temperature extraction of siderite in cold 10 % HCl for 24 hours, which simultaneously targets the ‘easily-reducible’ oxides (Table 5). Premature removal of ferrihydrite and lepidocrocite does not restrict the quantitative determination of Fe minerals in ancient sediment studies since the poorly-ordered, nanoparticulate oxyhydroxide minerals are rapidly transformed into the more thermodynamically-stable Fe mineral forms of goethite or haematite; ferrihydrite and lepidocrocite are therefore not found in ancient sediments (Cornell and Schwertmann, 2003). However, siderite, ferrihydrite, goethite, magnetite and nontronite are all important

constituents of modern sediments and their accurate quantitative identification is crucial for understanding the diagenetic process that impact global Fe, C, P, and S cycling (Lal, 2008; Cooper et al., 2005; Cooper et al., 2000; Kostka and Luther, 1994), the speciation and mobility of metals that contaminate natural environments (Jonsson and Sherman, 2008; Guo et al., 2007; Fendorf et al., 2000; Fredrickson et al., 2000; Lovley, 1993), and even the storage of radioactive waste (Hu et al., 2019; Ithurbide et al., 2010; Scheinost and Charlet, 2008).

The strength of any sequential extraction technique depends on the specific solubility of a particular mineral phase in a well-defined reagent. Incomplete, premature, and non-selective dissolution of target minerals exemplifies the importance of using an additional analytical technique to identify the residual phases at each stage of the procedure. The reported specificity of the sequential Fe extraction (Poulton and Canfield, 2005) cannot be assumed for all samples. When used in isolation, without careful calibration of the reacting Fe mineral phases, the sequential extraction could present entirely misleading information for the determination of specific Fe-containing minerals. Verification of the extracted or residual phases is therefore crucial for the correct interpretation of the leachate results.

Examples of secondary analyses used to verify the Fe minerals or mineral groups present in the extracted leachate or the residual material, include MBS (Jilbert et al., 2018; Sun et al., 2018; Schröder et al., 2016), LA-ICP-MS (Oonk et al., 2018), SEM or TEM (Xie et al., 2018; Zhuang et al., 2012), XAS (Sun et al., 2018; Barber et al., 2017), tests with  $^{58}\text{Fe}$  isotope spiking (Henkel et al. 2016), or a percentage yield comparison from a single mineral control or internal laboratory standard (Jin et al., 2018; Liu et al., 2018; Lynch et al., 2018; Qi et al., 2018; Lei et al., 2017; Oonk et al., 2017; Xie et al., 2017; Raiswell et al., 2010). The advantage of MBS is the ability to analyse a sample non-destructively and in a single, short step without additional preparation. XRD offers another suitable alternative for the

mineralogical characterisation of reactive Fe minerals when Co radiation (Mos et al., 2018) or a monochromator (Fransen, 2004) is used to repress fluorescence. However, X-ray diffractometers are rarely suitably configured for the classification of reactive Fe minerals. Although some of the verification techniques that are listed here are costly and not always readily available, there is little excuse not to run appropriate pure mineral standards alongside the extracted samples.

#### **4.2 Implications for use with modern and ancient sediments**

The implications of using data from unverified extractions, depend entirely upon the way in which it is interpreted. In ancient rocks and sediments that are diagenetically stabilised, the Poulton and Canfield (2005) chemical extraction is simply applied to compare the sum of highly reactive Fe (Fe-bound carbonates, oxides, magnetite, and pyrite) with that of total Fe, to distinguish oxic/euxinic from anoxic depositional conditions (Poulton et al., 2004). In fact, the Poulton and Canfield (2005) technique was specifically designed for this purpose.

Although the results of our study highlight some severe reliability issues with mineral specificity at individual stages, palaeoceanographic reconstructions of the water column only attempt to quantify the fraction of sediment Fe that is reactive towards sulphide (Berner, 1984, 1970). Since a mineral specific interpretation is not required, the non-specific removal of individual Fe phases observed at specific stages of a sequential extraction is unlikely to alter the conclusions of work that employs sediment Fe as a palaeoredox proxy. Furthermore, in many such studies, Fe palaeoredox data are used in combination with other palaeoenvironmental proxies to develop a picture of the evolving system.

Isotopic or trace metal analyses alternatively require the successful isolation of a specific mineral phase (e.g. carbonate), rather than an operationally-defined phase (e.g. Na acetate-extractable). Here, the Fe content of the leachate may only represent a small portion of the target mineral phase. As a purely hypothetical example, if less than half of the goethite and

other reducible iron oxides/oxyhydroxides are actually targeted during the  $\text{Fe}_{\text{di-ct}}$  reduction (Table 2 and Table 3), the estimated preservation of  $19 - 45 \times 10^{15}$  g of OC in surface marine sediments (Lalonde et al., 2012), and the impact of Fe in global C sequestration, would be severely underestimated. Studies based solely on the chemical extraction of Fe in modern sediments that assume a mineral-specific approach are most likely to be affected by inconsistent or erroneous outcomes of leaching methods.

Our work reiterates many of the limitations that have already been raised regarding the sequential extraction method for Fe (Poulton and Canfield, 2005). The incomplete extraction of siderite and goethite were respectively reported by Reinhard et al. (2009) and Claff et al. (2010). Both Schröder et al. (2016) and Henkel et al. (2016) documented the premature removal of magnetite, while Raiswell et al. (1994) demonstrated the extensive dissolution of nontronite prior to  $\text{Fe}_{\text{HCl}}$ . Our observation of premature ferrihydrite extraction during  $\text{Fe}_{\text{aca-48}}$  was also not surprising, since the 48-hour Na acetate leach at 50 °C is rarely used to extract siderite in samples where ferrihydrite is present. Furthermore, Poulton and Canfield (2005) reported the premature removal of small amounts of ferrihydrite (during the 24-hour  $\text{Fe}_{\text{aca}}$  stage) and magnetite (during the  $\text{Fe}_{\text{di-ct}}$  stage) in their original paper. In spite of such warnings, the Poulton and Canfield (2005) method continues to be used for mineral-specific analyses – a purpose for which it was not originally designed. Where sequential extractions are used to define specific mineral phases, we strongly advise that i) all leachate results are verified by a secondary, independent means of analysis and ii) the operational nature of extraction techniques is clearly defined.

## 5. CONCLUSIONS



The novel comparison of MBS residue and ICP-AAS leachate analyses at each stage in the Poulton and Canfield (2005) procedure, raises key concerns as to the reliable use of sequential extraction protocols for Fe mineral identification in mineral-specific applications. Our analyses use synthetic minerals produced according to Cornell and Schwertmann (2003), as in the original Poulton and Canfield (2005) method. Incomplete and premature dissolution of target minerals throughout the procedure affects the subsequent extraction stages. Unverified interpretations of the leachate Fe content are particularly misleading where the solubilised Fe is not the intended target phase. Matrix effects and grain size greatly affect the precision of individual extraction stages between natural and synthetic forms of the same mineral, between different natural samples of the same mineral, and even between identical samples. In fact, the behaviour of Fe minerals in the extraction method is clearly more dependent on the property of the mineral than the specific mineral itself. Poor precision and reliability are ultimately due to the operational nature of the technique, which is misunderstood or ignored in the majority of the research that employs the extraction for mineral-specific purposes.

MBS is a powerful tool with advantages over x-ray-based techniques, most notably the accurate characterisation of amorphous colloidal and nanoparticulate Fe minerals. The combined use of MBS alongside conventional sequential extraction procedures can provide a reliable means of semi-quantitative Fe mineral identification that is suitable for mineral specific applications in studies of both modern and ancient sediments.

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## REFERENCES

- Adamo, P., Agrelli, D. and Zampella, M. (2018) Chapter 9 - chemical speciation to assess bioavailability, bioaccessibility and geochemical forms of potentially toxic metals (PTMs) in polluted soils. In *Environmental Geochemistry*, second ed. (eds. B. De Vivo, H. E. Belkin, A. Lima). Elsevier (2018). pp 153-194.  
<http://dx.doi.org/10.1016/B978-0-444-63763-5.00010-0>.
- Afonso M. D. and Stumm W. (1992) Reductive dissolution of iron(III) (hydr)oxides by hydrogen sulfide. *Langmuir* **8**, 1671-1675. <https://doi.org/10.1021/la00042a030>.
- Bacon J. R. and Davidson C. M. (2008) Is there a future for sequential chemical extraction? *Analyst* **133**, 25-46. <https://doi.org/10.1039/B711896A>.
- Baeyens W., Monteny F., Leermakers M. and Bouillon S. (2003) Evaluation of sequential extractions on dry and wet sediments. *Anal. Bioanal. Chem.* **376**, 890-901.  
<https://doi.org/10.1007/s00216-003-2005-z>.
- Barber A., Brandes J., Leri A., Lalonde K., Balind K., Wirick S., Wang J. and Gélinas Y. (2017) Preservation of organic matter in marine sediments by inner-sphere interactions with reactive iron. *Sci. Rep.-UK* **7**, 366. <https://doi.org/10.1038/s41598-017-00494-0>.
- Berner R. A. (1970) Sedimentary pyrite formation. *Am. J. Sci.* **268**, 1-23.  
<https://doi.org/10.2475/ajs.268.1.1>.

- Berner R. A. (1984) Sedimentary pyrite formation - an update. *Geochim. Cosmochim. Acta* **48**, 605-615. [https://doi.org/10.1016/0016-7037\(84\)90089-9](https://doi.org/10.1016/0016-7037(84)90089-9).
- Blonder B., Boyko V., Turchyn A.V., Antler G., Sinichkin U., Knossow N., Klein R. and Kamyshny A. (2017) Impact of aeolian dry deposition of reactive iron minerals on sulfur cycling in sediments of the Gulf of Aqaba. *Front. Microbiol.* **8**, 1131. <https://doi.org/10.3389/fmicb.2017.01131>.
- Borggaard O.K. (1988) Phase identification by phase dissolution techniques. In *Iron in soils and clay minerals* NATO ASI Series (Series C: Mathematical and Physical Sciences), vol 217 (eds. J. W. Stucki, B. A. Goodman, U. Schwertmann). Springer, Dordrecht. pp. 83-98. [https://doi.org/10.1007/978-94-009-4007-9\\_5](https://doi.org/10.1007/978-94-009-4007-9_5).
- Brindley G. W. (1980) Order-disorder in clay mineral structures. In *Crystal structures of clay minerals and their x-ray identification* (eds G. W. Brindley and G. Brown). Mineralogical Society, London. pp. 125-195. <https://doi.org/10.1180/mono-5.2>.
- Canfield D. E. (1989) Reactive iron in marine sediments. *Geochim Cosmochim Acta* **53**, 619-632. [https://doi.org/10.1016/0016-7037\(89\)90005-7](https://doi.org/10.1016/0016-7037(89)90005-7).
- Canfield D. E., Poulton S. W. and Narbonne G. M. (2007) Late-Neoproterozoic deep-ocean oxygenation and the rise of animal life. *Science* **315**, 92-95. <https://doi.org/10.1126/science.1135013>.
- Chen X., Ling H. F., Vance D., Shields-Zhou G. A., Zhu M. Y., Poulton S. W., Och L. M., Jiang S. Y., Li D., Cremonese L. and Archer C. (2015) Rise to modern levels of ocean oxygenation coincided with the Cambrian radiation of animals. *Nat. Commun.* **6**, 7142. <https://doi.org/10.1038/ncomms8142>.
- Claff S. R., Sullivan L. A., Burton E. D. and Bush R. T. (2010) A sequential extraction procedure for acid sulfate soils: partitioning of iron. *Geoderma* **155**, 224-230. <https://doi.org/10.1016/j.geoderma.2009.12.002>.

- Cooper D. C., Neal A. L., Kukkadapu R. K., Brewe D., Coby A. and Picardal F. W. (2005) Effects of sediment iron mineral composition on microbially mediated changes in divalent metal speciation: importance of ferrihydrite. *Geochim. Cosmochim. Acta* **69**, 1739-1754. <https://doi.org/10.1016/j.gca.2004.09.013>.
- Cooper D. C., Picardal F., Rivera J. and Talbot C. (2000) Zinc immobilization and magnetite formation via ferric oxide reduction by *Shewanella putrefaciens* 200. *Environ. Sci. Technol.* **34**, 100-106. <https://doi.org/10.1021/es990510x>.
- Cornell R. M. and Schwertmann U. (2003) Synthesis. In *The iron oxides: structure, properties, reactions, occurrences and uses* (eds. R. M. Cornell and U. Schwertmann). WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim. pp. 527-542. <https://doi.org/10.1002/3527602097.ch20>.
- Dahl T. W., Hammarlund E. U., Anbar A. D., Bond D. P. G., Gill B. C., Gordon G. W., Knoll A. H., Nielsen A. T., Schovsbo N. H. and Canfield D. E. (2010) Devonian rise in atmospheric oxygen correlated to the radiations of terrestrial plants and large predatory fish. *P. Natl. Acad. Sci. USA* **107**, 17911-17915. <https://doi.org/10.1073/pnas.1011287107>.
- Dijkstra N., Slomp C. P., Behrends T. and Expedition 347 Scientists (2016) Vivianite is a key sink for phosphorus in sediments of the Landsort Deep, an intermittently anoxic deep basin in the Baltic Sea. *Chem. Geol.* **438**, 58-72. <https://doi.org/10.1016/j.chemgeo.2016.05.025>.
- Ding S., Wang Y., Wang D., Li Y. Y., Gong M. and Zhang C. (2016) In situ, high-resolution evidence for iron-coupled mobilization of phosphorus in sediments. *Sci. Rep.-UK* **6**, 24341. <https://doi.org/10.1038/srep24341>.

- Eiche E., Kramar U., Berg M., Berner Z., Norra S. and Neumann T. (2010) Geochemical changes in individual sediment grains during sequential arsenic extractions. *Water Res.* **44**, 5545-5555. <https://doi.org/10.1016/j.watres.2010.06.002>.
- Estes E. R., Pockalny R., D'Hondt S., Inagaki F., Morono Y., Murray R. W., Nordlund D., Spivack A. J., Wankel S. D., Xiao N. and Hansel C. M. (2019) Persistent organic matter in oxic subseafloor sediment. *Nat. Geosci.* **12**, 126-131. <https://doi.org/10.1038/s41561-018-0291-5>.
- Fendorf S., Wielinga B. W. and Hansel C. M. (2000) Chromium transformations in natural environments: the role of biological and abiological processes in chromium(VI) reduction. *Int. Geol. Rev.* **42**, 691-701. <https://doi.org/10.1080/00206810009465107>.
- Fialips C. I., Huo D. F., Yan L. B., Wu J. and Stucki J. W. (2002) Effect of Fe oxidation state on the IR spectra of Garfield nontronite. *Am. Mineral.* **87**, 630-641. <https://doi.org/10.2138/am-2002-5-605>.
- Fransen, M. J. (2004) 1- and 2-dimensional detection systems and the problem of sample fluorescence in x-ray diffractometry. *Adv X-ray Anal.* **47**, 224–231.
- Fredrickson J. K., Zachara J. M., Kennedy D. W., Duff M. C., Gorby Y. A., Li S.M.W. and Krupka K. M. (2000) Reduction of U(VI) in goethite ( $\alpha$ -FeOOH) suspensions by a dissimilatory metal-reducing bacterium. *Geochim. Cosmochim. Acta* **64**, 3085-3098. [https://doi.org/10.1016/S0016-7037\(00\)00397-5](https://doi.org/10.1016/S0016-7037(00)00397-5).
- Ghaisas N. A., Maiti K. and White J. R. (2019) Coupled iron and phosphorus release from seasonally hypoxic Louisiana shelf sediment. *Estuar. Coast. Shelf S.* **219**, 81-89. <https://doi.org/10.1016/j.ecss.2019.01.019>.
- Gobeil C., Johnson W. K., Macdonald R. W. and Wong C. S. (1995) Sources and burden of lead in St. Lawrence estuary sediments – isotopic evidence. *Environ. Sci. & Technol.* **29**, 193-201. <https://doi.org/10.1021/es00001a025>.

- Gómez-Ariza, J. L., Giráldez, I., Sánchez-Rodas, D. and Morales, E. (1999) Metal readsorption and redistribution during the analytical fractionation of trace elements in oxic estuarine sediments. *Anal. Chim. Acta* **399**, 295-307.  
[https://doi.org/10.1016/S0003-2670\(99\)00460-2](https://doi.org/10.1016/S0003-2670(99)00460-2).
- Guilbaud R., Butler I. B., Ellam R. M. and Rickard, D. (2010) Fe isotope exchange between  $\text{Fe(II)}_{\text{aq}}$  and nanoparticulate mackinawite ( $\text{FeS}_m$ ) during nanoparticle growth. *Earth Planet. Sc. Lett.* **300**, 174-183. <https://doi.org/10.1016/j.epsl.2010.10.004>.
- Guo H. M., Stüben D. and Berner Z. (2007) Removal of arsenic from aqueous solution by natural siderite and hematite. *Appl. Geochem.* **22**, 1039-1051.  
<https://doi.org/10.1016/j.apgeochem.2007.01.004>.
- Gütlich P. and Schröder C. (2012) Mössbauer spectroscopy. In *Methods in Physical Chemistry* (eds. R. Schäfer and P. C. Schmidt). Wiley-VCH, Weinheim. pp. 351-389.  
<https://doi.org/10.1002/9783527636839.ch11>.
- Haese R. R., Wallmann K., Dahmke A., Kretzmann U., Müller P. J. and Schulz H.D. (1997) Iron species determination to investigate early diagenetic reactivity in marine sediments. *Geochim. Cosmochim. Acta* **61**, 63-72. [https://doi.org/10.1016/S0016-7037\(96\)00312-2](https://doi.org/10.1016/S0016-7037(96)00312-2).
- Hammarlund E. U., Smith M. P., Rasmussen J. A., Nielsen A. T., Canfield D. E. and Harper D. A. T. (2019) The Sirius Passet Lagerstätte of North Greenland—geochemical window on early Cambrian low-oxygen environments and ecosystems. *Geobiology* **17**, 12-26. <https://doi.org/10.1111/gbi.12315>.
- Hanahan C. (2004) Dissolution of hydroxide minerals in the 1 M sodium acetate, pH 5, extracting solution in sequential extraction schemes. *Environ. Geol.* **45**, 864-868.  
<https://doi.org/10.1007/s00254-003-0946-3>.

- Hass A. and Fine P. (2010) Sequential selective extraction procedures for the study of heavy metals in soils, sediments, and waste materials—a critical review. *Crit. Rev. Env. Sci. Tec.* **40**, 365-399. <https://doi.org/10.1080/10643380802377992>.
- Henkel S., Kasten S., Poulton S. W. and Staubwasser M. (2016) Determination of the stable iron isotopic composition of sequentially leached iron phases in marine sediments. *Chem. Geol.* **421**, 93-102. <https://doi.org/10.1016/j.chemgeo.2015.12.003>.
- Henkel S., Kasten S., Hartmann J. F., Silva-Busso A. and Staubwasser M. (2018) Iron cycling and stable Fe isotope fractionation in Antarctic shelf sediments, King George Island. *Geochim. Cosmochim. Acta* **237**, 320-338. <https://doi.org/10.1016/j.gca.2018.06.042>.
- Heron G., Crouzet C., Bourg A. C. M. and Christensen T. H. (1994) Speciation of Fe(II) and Fe(III) in contaminated aquifer sediments using chemical extraction techniques. *Environ. Sci. Tech.* **28**, 1698-1705. <https://doi.org/10.1021/es00058a023>.
- Homoky W. B., John S. G., Conway T. M. and Mills R. A. (2013) Distinct iron isotopic signatures and supply from marine sediment dissolution. *Nat. Commun.* **4**, 2143. <https://doi.org/10.1038/ncomms3143>.
- Hu W., Zhang Z. X., Li M. X., Liu H. B., Zhang C. G., Chen T. H. and Zhou Y. F. (2019) Enhanced uptake capacity for uranium(VI) in aqueous solutions by activated natural siderite: performance and mechanism. *Appl. Geochem.* **100**, 96-103. <https://doi.org/10.1016/j.apgeochem.2018.11.010>.
- Ithurbide A., Peulon S., Miserque F., Beaucaire C. and Chaussé, A. (2010) Retention and redox behaviour of uranium(VI) by siderite (FeCO<sub>3</sub>). *Radiochim. Acta* **98**, 563-568. <https://doi.org/10.1524/ract.2010.1754>.

- Jaisi D. P., Dong H. L. and Morton J. P. (2008) Partitioning of Fe(II) in reduced nontronite (NAu-2) to reactive sites: reactivity in terms of Tc(VII) reduction. *Clay. Clay Miner.* **56**, 175-189. <https://doi.org/10.1346/CCMN.2008.0560204>.
- Jilbert T., Asmala E., Schröder C., Tiihonen R., Myllykangas J. P., Virtasalo J. J., Kotilainen A., Peltola P., Ekholm P. and Hietanen S. (2018) Impacts of flocculation on the distribution and diagenesis of iron in boreal estuarine sediments. *Biogeosciences* **15**, 1243-1271. <https://doi.org/10.5194/bg-15-1243-2018>.
- Jin C. S., Li C., Algeo T. J., O'Connell B., Cheng M., Shi W., Shen J. and Planavsky N. J. (2018) Highly heterogeneous "poikiloredox" conditions in the early Ediacaran Yangtze Sea. *Precambrian Res.* **311**, 157-166. <https://doi.org/10.1016/j.precamres.2018.04.012>.
- Jönsson J. and Sherman D. M. (2008) Sorption of As(III) and As(V) to siderite, green rust (fougerite) and magnetite: implications for arsenic release in anoxic groundwaters. *Chem. Geol.* **255**, 173-181. <https://doi.org/10.1016/j.chemgeo.2008.06.036>.
- Keil R. G., Montlucon D. B., Prah F. G. and Hedges J. I. (1994) Sorptive preservation of labile organic matter in marine sediments. *Nature* **370**, 549-552. <https://doi.org/10.1038/370549a0>.
- Kersten M., Garbe-Schönberg C. D., Thomsen S., Anagnostou C. and Sioulas A. (1997) Source apportionment of Pb pollution in the coastal waters of Elefsis Bay, Greece. *Environ. Sci. Technol.* **31**, 1295-1301. <https://doi.org/10.1021/es960473z>.
- Kheboian C. and Bauer C. F. (1987) Accuracy of selective extraction procedures for metal speciation in model aquatic sediments. *Anal. Chem.* **59**, 1417-1423. <https://doi.org/10.1021/ac00137a010>.
- Kim E. J., Lee J. C. and Baek K. (2015) Abiotic reductive extraction of arsenic from contaminated soils enhanced by complexation: arsenic extraction by reducing agents



- and combination of reducing and chelating agents. *J. Hazard. Mater.* **283**, 454-461.  
<https://doi.org/10.1016/j.jhazmat.2014.09.055>.
- Kostka J. E. and Luther III G. W. (1994) Partitioning and speciation of solid-phase iron in salt marsh sediments. *Geochim. Cosmochim. Acta* **58**, 1701-1710.  
[https://doi.org/10.1016/0016-7037\(94\)90531-2](https://doi.org/10.1016/0016-7037(94)90531-2).
- Kraal P., Dijkstra N., Behrends T. and Slomp C. P. (2017) Phosphorus burial in sediments of the sulfidic deep Black Sea: key roles for adsorption by calcium carbonate and apatite authigenesis. *Geochim. Cosmochim. Acta* **204**, 140-158.  
<https://doi.org/10.1016/j.gca.2017.01.042>.
- Lal R. (2008) Sequestration of atmospheric CO<sub>2</sub> in global carbon pools. *Energ. Environ. Sci.* **1**, 86-100. <https://doi.org/10.1039/B809492F>.
- Lalonde K., Mucci A., Ouellet A. and G  linas Y. (2012) Preservation of organic matter in sediments promoted by iron. *Nature* **483**, 198-200.  
<https://doi.org/10.1038/nature10855>.
- Lei L.-D., Shen J., Li C., Algeo T. J., Chen Z.-Q., Feng Q.-L., Cheng M., Jin C.-S. and Huang J.-H. (2017) Controls on regional marine redox evolution during Permian-Triassic transition in South China. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* **486**, 17-32. <https://doi.org/10.1016/j.palaeo.2017.02.010>.
- Lenstra W. K., Egger M., van Helmond N. A. G. M., Kritzberg E., Conley D. J. and Slomp C. P. (2018) Variations in iron input to an oligotrophic Baltic Sea estuary: impact on sedimentary phosphorus burial. *Biogeosciences* **15**, 6979-6996.  
<https://doi.org/10.5194/bg-15-6979-2018>.
- Li J., Zhang Y. and Katsev S. (2018) Phosphorus recycling in deeply oxygenated sediments in Lake Superior controlled by organic matter mineralization. *Limnol. Oceanogr.* **63**, 1372-1385. <https://doi.org/10.1002/lno.10778>.

- Liu Y., Zhang J. and He H. (2018) Assessment of the Tessier and BCR sequential extraction procedures for elemental partitioning of Ca, Fe, Mn, Al, and Ti and their application to surface sediments from Chinese continental shelf. *Acta Oceanol. Sin.* **37**, 22-28. <https://doi.org/10.1007/s13131-018-1189-1>.
- Lovley D. R. (1993) Dissilatory metal reduction. *Annu. Rev. Microbiol.* **47**, 263-290. [10.1146/annurev.mi.47.100193.001403](https://doi.org/10.1146/annurev.mi.47.100193.001403).
- Lynch S. F. L., Batty L. C. and Byrne P. (2018) Environmental risk of severely Pb-contaminated riverbank sediment as a consequence of hydrometeorological perturbation. *Sci. Total Environ.* **636**, 1428-1441. <https://doi.org/10.1016/j.scitotenv.2018.04.368>.
- Mos Y. M., Vermeulen A. C., Buisman C. J. N. and Weijma J. (2018) X-ray diffraction of iron containing samples: the importance of a suitable configuration. *Geomicrobiol. J.* **35**, 511-517. <https://doi.org/10.1080/01490451.2017.1401183>.
- Ma W.-W., Zhu M.-X., Yang G.-P. and Li, T. (2017) In situ, high-resolution DGT measurements of dissolved sulfide, iron and phosphorus in sediments of the East China Sea: insights into phosphorus mobilization and microbial iron reduction. *Mar. Pollut. Bull.* **124**, 400-410. <https://doi.org/10.1016/j.marpolbul.2017.07.056>.
- Ma W.-W., Zhu M.-X., Yang G.-P. and Li T. (2018) Iron geochemistry and organic carbon preservation by iron (oxyhydr)oxides in surface sediments of the East China Sea and the south Yellow Sea. *J. Marine Syst.* **178**, 62-74. <https://doi.org/10.1016/j.jmarsys.2017.10.009>.
- Manceau A., Drits V. A., Lanson B., Chateigner D., Wu J., Huo D., Gates W. P. and Stucki J. W. (2000) Oxidation-reduction mechanism of iron in dioctahedral smectites: II. Crystal chemistry of reduced Garfield nontronite. *Am. Mineral.* **85**, 153-172. <https://doi.org/10.2138/am-2000-0115>.

- Markussen T. N., Elberling B., Winter C. and Andersen T. J. (2016) Flocculated meltwater particles control Arctic land-sea fluxes of labile iron. *Sci. Rep.-UK* **6**, 24033. <https://doi.org/10.1038/srep24033>.
- Mayer L. M. (1994) Surface area control of organic carbon accumulation in continental shelf sediments. *Geochim. Cosmochim. Acta* **58**, 1271-1284. [https://doi.org/10.1016/0016-7037\(94\)90381-6](https://doi.org/10.1016/0016-7037(94)90381-6).
- Moreira-Chavarría A., Griffioen J. and Behrends T. (2016) Optimized sequential extraction for carbonates: quantification and  $\delta^{13}\text{C}$  analysis of calcite, dolomite and siderite. *Chem. Geol.* **443**, 146-157. <https://doi.org/10.1016/j.chemgeo.2016.09.025>.
- Nemati K., Bakar N. K. A., Abas M. R. and Sobhanzadeh E. (2009) Concentration measurement and evaluation of mobility of heavy metals of Zayandeh-Rood river sediments. *Asian J. Chem.* **21**, 4894-4900.
- Oonk P. B. H., Mason P. R. D., Tsikos H. and Bau M. (2018) Fraction-specific rare earth elements enable the reconstruction of primary seawater signatures from iron formations. *Geochim. Cosmochim. Acta* **238**, 102-122. <https://doi.org/10.1016/j.gca.2018.07.005>.
- Oonk P. B. H., Tsikos H., Mason P. R. D., Henkel S., Staubwasser M., Fryer L., Poulton S. W. and Williams H.M. (2017) Fraction-specific controls on the trace element distribution in iron formations: implications for trace metal stable isotope proxies. *Chem. Geol.* **474**, 17-32. <https://doi.org/10.1016/j.chemgeo.2017.10.018>.
- Parat C., Lévêque J., Dousset S., Chaussod R. and Andreux F. (2003) Comparison of three sequential extraction procedures used to study trace metal distribution in an acidic sandy soil. *Anal. Bioanal. Chem.* **376**, 243-247. <https://doi.org/10.1007/s00216-003-1864-7>.

- Peter S. and Sobek S. (2018) High variability in iron-bound organic carbon among five boreal lake sediments. *Biogeochemistry* **139**, 19-29. <https://doi.org/10.1007/s10533-018-0456-8>.
- Posth N. R., Köhler I., Swanner E. D., Schröder C., Wellmann E., Binder B., Konhauser K. O., Neumann U., Berthold C., Nowak M. and Kappler A. (2013) Simulating Precambrian banded iron formation diagenesis. *Chem. Geol.* **362**, 66-73. <https://doi.org/10.1016/j.chemgeo.2013.05.031>.
- Poulton S. W. and Canfield D. E. (2005) Development of a sequential extraction procedure for iron: implications for iron partitioning in continentally derived particulates. *Chem. Geol.* **214**, 209-221. <https://doi.org/10.1016/j.chemgeo.2004.09.003>.
- Poulton S. W., Fralick P. W. and Canfield D. E. (2004) The transition to a sulphidic ocean similar to 1.84 billion years ago. *Nature* **431**, 173-177. <https://doi.org/10.1038/nature02912>.
- Qi C., Li C., Gabbott S. E., Ma X., Xie L., Deng W., Jin C. and Hou X.-G. (2018) Influence of redox conditions on animal distribution and soft-bodied fossil preservation of the Lower Cambrian Chengjiang Biota. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* **507**, 180-187. <https://doi.org/10.1016/j.palaeo.2018.07.010>.
- Raiswell R., Benning L. G., Tranter M. and Tulaczyk S. (2008) Bioavailable iron in the Southern Ocean: the significance of the iceberg conveyor belt. *Geochem T.* **9**, 7. <https://doi.org/10.1186/1467-4866-9-7>.
- Raiswell R. and Canfield D. E. (1996) Rates of reaction between silicate iron and dissolved sulfide in Peru Margin sediments. *Geochim. Cosmochim. Acta* **60**, 2777-2787. [https://doi.org/10.1016/0016-7037\(96\)00141-X](https://doi.org/10.1016/0016-7037(96)00141-X).
- Raiswell R., Canfield D. E. and Berner R. A. (1994) A comparison of iron extraction methods for the determination of degree of pyritisation and the recognition of iron-

- limited pyrite formation. *Chem. Geol.* **111**, 101-110. [https://doi.org/10.1016/0009-2541\(94\)90084-1](https://doi.org/10.1016/0009-2541(94)90084-1).
- Raiswell R., Reinhard C. T., Derkowski A., Owens J., Bottrell S. H., Anbar A. D. and Lyons T. W. (2011) Formation of syngenetic and early diagenetic iron minerals in the late Archean Mt. McRae Shale, Hamersley Basin, Australia: new insights on the patterns, controls and paleoenvironmental implications of authigenic mineral formation. *Geochim. Cosmochim. Acta* **75**, 1072-1087. <https://doi.org/10.1016/j.gca.2010.11.013>.
- Raiswell R., Vu H. P., Brinza L. and Benning L. G. (2010) The determination of labile Fe in ferrihydrite by ascorbic acid extraction: methodology, dissolution kinetics and loss of solubility with age and de-watering. *Chem. Geol.* **278**, 70-79. <https://doi.org/10.1016/j.chemgeo.2010.09.002>.
- Rancourt D. G. and Ping J. Y. (1991) Voigt-based methods for arbitrary-shape static hyperfine parameter distributions in Mössbauer spectroscopy. *Nucl. Instrum. Meth. B* **58**, 85-97. [https://doi.org/10.1016/0168-583X\(91\)95681-3](https://doi.org/10.1016/0168-583X(91)95681-3).
- Ransom B., Kim D., Kastner M. and Wainwright S. (1998) Organic matter preservation on continental slopes: importance of mineralogy and surface area. *Geochim. Cosmochim. Acta* **62**, 1329-1345. [https://doi.org/10.1016/S0016-7037\(98\)00050-7](https://doi.org/10.1016/S0016-7037(98)00050-7).
- Reinhard C. T., Raiswell R., Scott C., Anbar A. D. and Lyons T. W. (2009) A late Archean sulfidic sea stimulated by early oxidative weathering of the continents. *Science* **326**, 713-716. <https://doi.org/10.1126/science.1176711>.
- Ryan P. C., Hillier S. and Wall A. J. (2008) Stepwise effects of the BCR sequential chemical extraction procedure on dissolution and metal release from common ferromagnesian clay minerals: a combined solution chemistry and X-ray powder diffraction study. *Sci. Total Environ.* **407**, 603-614. <https://doi.org/10.1016/j.scitotenv.2008.09.019>.

- Sahuquillo A., López-Sánchez, J. F., Rubio R., Rauret G., Thomas R. P., Davidson C. M. and Ure A. M. (1999) Use of a certified reference material for extractable trace metals to assess sources of uncertainty in the BCR three-stage sequential extraction procedure. *Anal. Chim. Acta* **382**, 317-327. [https://doi.org/10.1016/S0003-2670\(98\)00754-5](https://doi.org/10.1016/S0003-2670(98)00754-5).
- Sanchez-Espana J. and Reyes J. (2019) Comparing schwertmannite and hydrobasaluminite dissolution in ammonium oxalate (pH 3.0): implications for metal speciation studies by sequential extraction. *Minerals* **9**, 57. <https://doi.org/10.3390/min9010057>.
- Scheinost A. C. and Charlet L. (2008) Selenite reduction by mackinawite, magnetite and siderite: XAS characterization of nanosized redox products. *Environ. Sci. Technol.* **42**, 1984-1989. <https://doi.org/10.1021/es071573f>.
- Schindelin J., Arganda-Carreras I., Frise E., Kaynig V., Longair M., Pietzsch T., Preibisch S., Preibisch S., Rueden C., Saalfeld S., Schmid B., Tinevez J. Y., White D. J., Hartenstein V., Eliceiri K., Tomancak P. and Cardona A. (2012) Fiji: an open source platform for biological-image analysis. *Nat. Methods* **9**, 676 - 682. <https://doi.org/10.1038/nmeth.2019>.
- Scholz F., Severmann S., McManus J., Noffke A., Lomnitz U. and Hensen C. (2014) On the isotope composition of reactive iron in marine sediments: redox shuttle versus early diagenesis. *Chem. Geol.* **389**, 48-59. <https://doi.org/10.1016/j.chemgeo.2014.09.009>.
- Schröder C., Köhler I., Muller, F. L. L., Chumakov A. I., Kuppenko I., Rüffer R. and Kappler A. (2016) The biogeochemical iron cycle and astrobiology. *Hyperfine Interact.* **237**, 85. <https://doi.org/10.1007/s10751-016-1289-2>.
- Severmann S., McManus J., Berelson W. M. and Hammond D. E. (2010) The continental shelf benthic iron flux and its isotope composition. *Geochim. Cosmochim. Acta* **74**, 3984-4004. <https://doi.org/10.1016/j.gca.2010.04.022>.

- Sulkowski M. and Hirner A. V. (2006) Element fractionation by sequential extraction in a soil with high carbonate content. *Appl. Geochem.* **21**, 16-28.  
<https://doi.org/10.1016/j.apgeochem.2005.09.016>.
- Sun J., Mailloux B. J., Chillrud S. N., van Geen A., Thompson A. and Bostick B. C. (2018) Simultaneously quantifying ferrihydrite and goethite in natural sediments using the method of standard additions with X-ray absorption spectroscopy. *Chem. Geol.* **476**, 248-259. <https://doi.org/10.1016/j.chemgeo.2017.11.021>.
- Sutherland R. A. (2010) BCR (R)-701: a review of 10-years of sequential extraction analyses. *Anal. Chim. Acta* **680**, 10-20. <https://doi.org/10.1016/j.aca.2010.09.016>.
- Thompson J., Poulton S. W., Guilbaud R., Doyle K. A., Reid S. and Krom M.D. (2019). Development of a modified SEDEX phosphorus speciation method for ancient rocks and modern iron-rich sediments. *Chem. Geol.* **524**, 383-393.  
<https://doi.org/10.1016/j.chemgeo.2019.07.003>
- Tlustoš P., Száková J., Stárková A. and Pavlíková D. (2005) A comparison of sequential extraction procedures for fractionation of arsenic, cadmium, lead, and zinc in soil. *Cent. Eur. J. Chem.* **3**, 830-851. <https://doi.org/10.2478/BF02475207>.
- Ure A. M. (1991) Trace element speciation in soils, soil extracts and solutions. *Microchim. Acta* **104**, 49. <https://doi.org/10.1007/BF01245495>.
- Wallmann K., Hennies K., König I., Petersen, W. and Knauth H. D. (1993) New procedure for determining reactive Fe(III) and Fe(II) minerals in sediments. *Limnol. Oceanogr.* **38**, 1803-1812. <https://doi.org/10.4319/lo.1993.38.8.1803>.
- Weber F.-A., Voegelin A. and Kretzschmar R. (2009) Multi-metal contaminant dynamics in temporarily flooded soil under sulfate limitation. *Geochim. Cosmochim. Acta* **73**, 5513-5527. <https://doi.org/10.1016/j.gca.2009.06.011>.

- Wehrmann L. M., Riedinger N., Brunner B., Kamyshny Jr. A., Hubert C. R. J., Herbert L. C., Brüchert, V., Jørgensen B. B., Ferdelman T. G. and Formolo M. J. (2017) Iron-controlled oxidative sulfur cycling recorded in the distribution and isotopic composition of sulfur species in glacially influenced fjord sediments of west Svalbard. *Chem. Geol.* **466**, 678-695. <https://doi.org/10.1016/j.chemgeo.2017.06.013>.
- Whalley C. and Grant, A. (1994) Assessment of the phase selectivity of the European Community Bureau of Reference (BCR) sequential extraction procedure for metals in sediment. *Anal. Chim. Acta* **291**, 287-295. [https://doi.org/10.1016/0003-2670\(94\)80024-3](https://doi.org/10.1016/0003-2670(94)80024-3).
- Wu T., Shelobolina E., Xu H. F., Konishi H., Kukkadapu R. and Roden E. E. (2012) Isolation and microbial reduction of Fe(III) phyllosilicates from subsurface sediments. *Environ. Sci. Technol.* **46**, 11618-11626. <https://doi.org/10.1021/es302639n>.
- Xie Y., Lu G., Yang C., Qu L., Chen M., Guo C. and Dang Z. (2018) Mineralogical characteristics of sediments and heavy metal mobilization along a river watershed affected by acid mine drainage. *Plos One* **13**, e0190010. <https://doi.org/10.1371/journal.pone.0190010>.
- Xie Y., Lu G., Ye H., Yang C., Xia D., Yi X., Reinfelder J. and Dang Z. (2017) Fulvic acid induced the liberation of chromium from  $\text{CrO}_4^{2-}$ -substituted schwertmannite. *Chem. Geol.* **475**, 52-61. <https://doi.org/10.1016/j.chemgeo.2017.10.031>.
- Zhuang Y.-F., Fialips C. I., White M. L. and Ferrandez D. M. P. (2012) New redox-active material for permeable water remediation systems. *Appl. Clay Sci.* **59-60**, 26-35. <https://doi.org/10.1016/j.clay.2012.02.002>.
- Zimmerman, A. J. and Weindorf, D. C. (2010) Heavy metal and trace metal analysis in soil by sequential extraction: a review of procedures. *Int. J. Anal. Chem.* **2010**, 387803. <https://doi.org/10.1155/2010/387803>.



Appendix A to

**The use of operationally-defined sequential Fe extraction methods for mineralogical applications: a cautionary tale from Mössbauer spectroscopy**

*Laura E. Hepburn, Ian B. Butler, Adrian Boyce, Christian Schröder*

**Content**

**Figure A1:** Unleached natural Ivigtut siderite XRD spectrum.

**Figure A2:** Unleached synthetic 2-line ferrihydrite XRD spectrum.

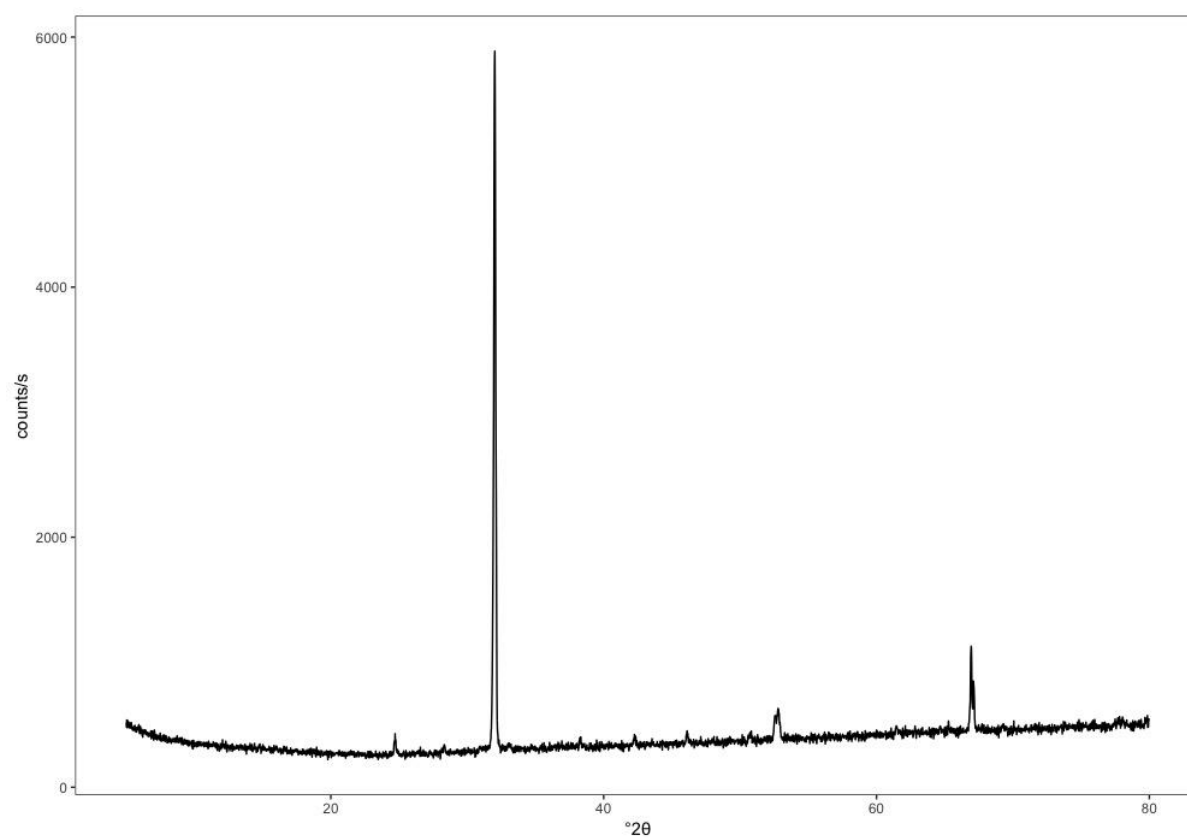
**Figure A3:** Unleached synthetic goethite XRD spectrum.

**Figure A4:** Unleached natural UoE goethite XRD spectrum.

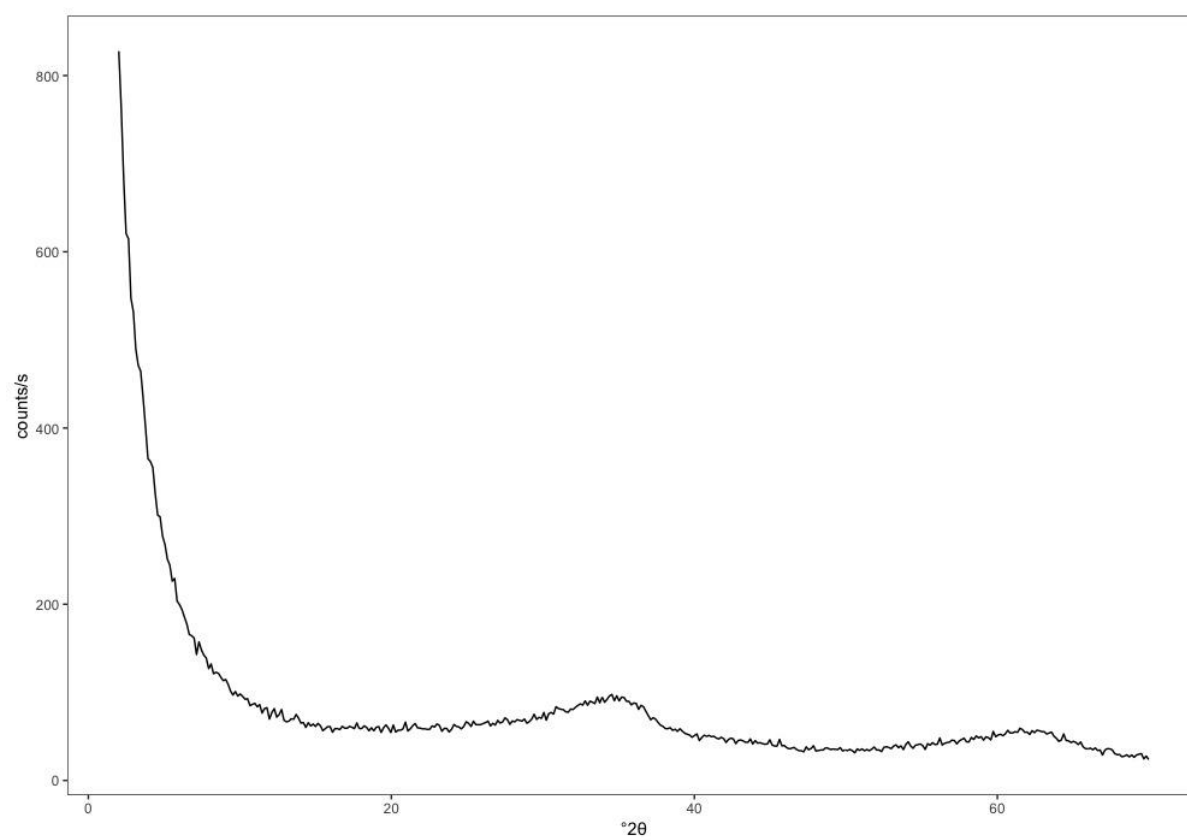
**Figure A5:** Unleached synthetic magnetite XRD spectrum.

**Figure A6:** Unleached natural NMS nontronite XRD spectrum.

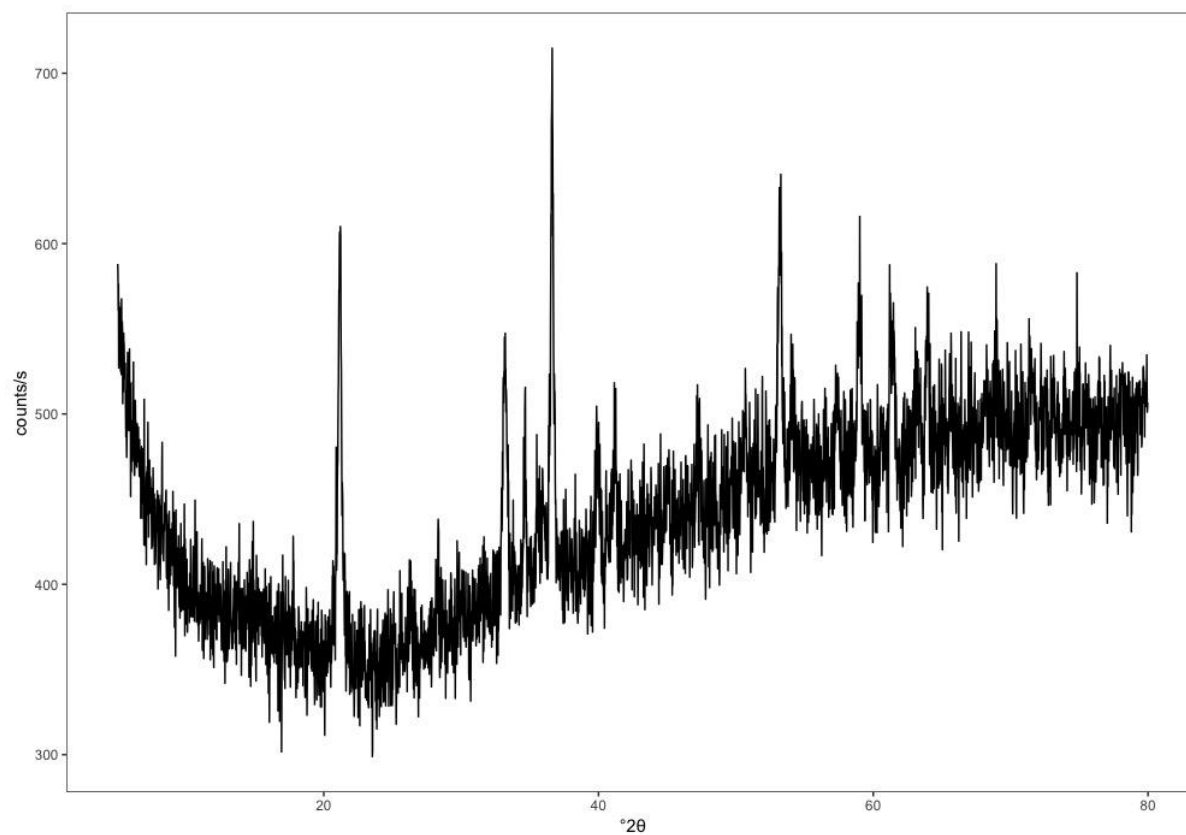
**Figure A7:** Unleached natural NAu-2 nontronite XRD spectrum.



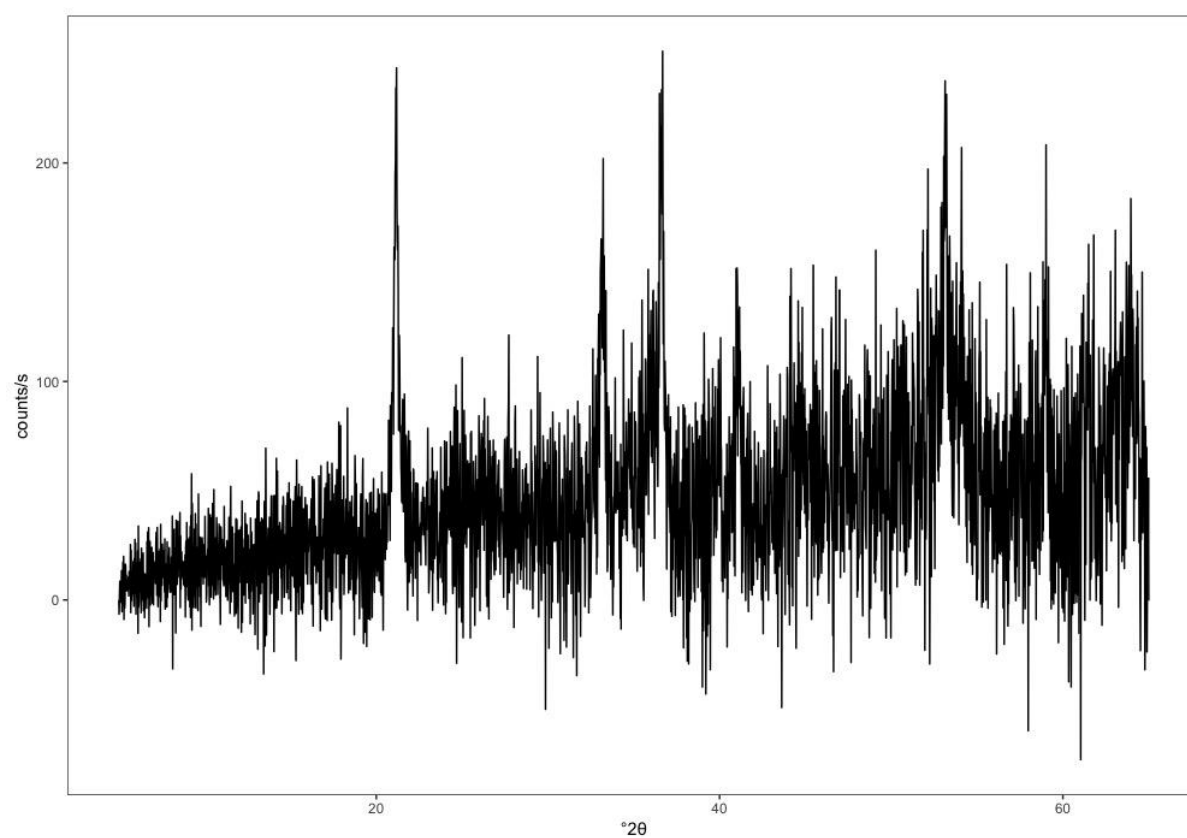
*Figure A1: Unleached natural Ivigtut siderite XRD spectrum. CuK $\alpha$  radiation without monochromator, National Museums Collection Centre, Edinburgh, UK.*



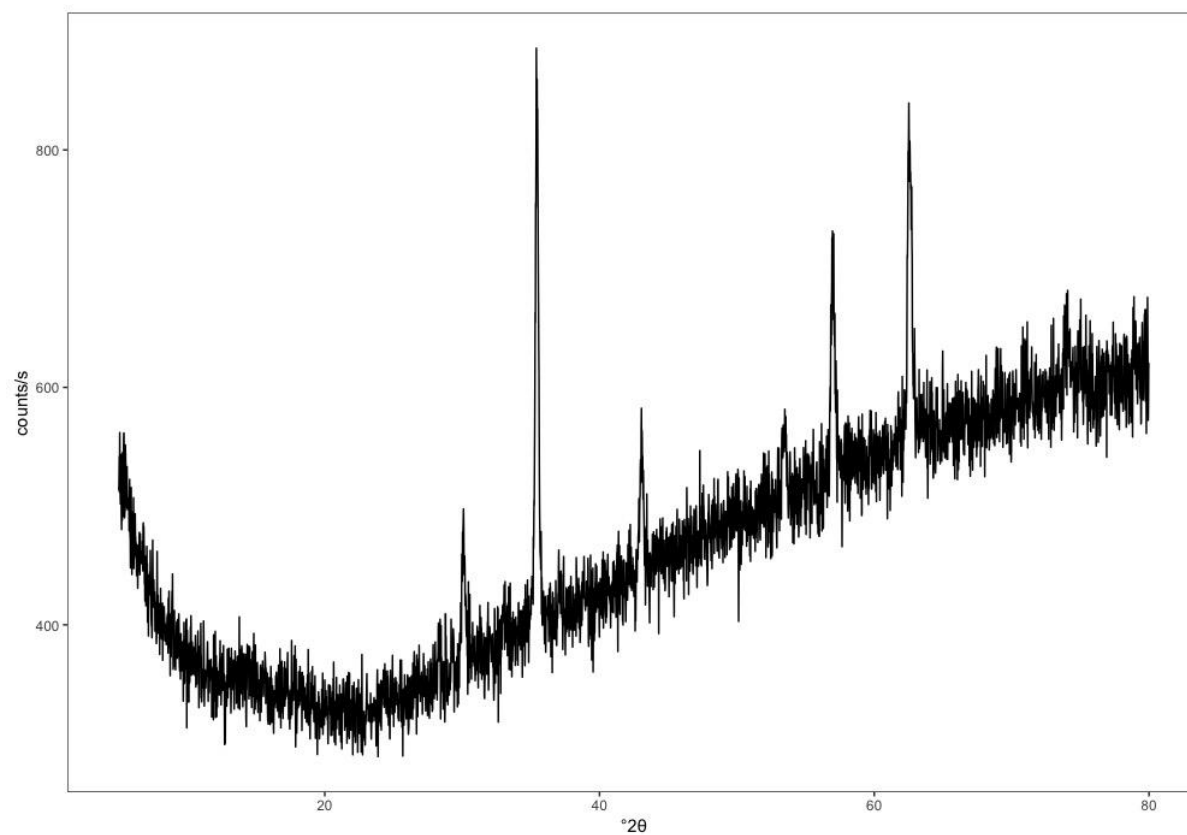
*Figure A2: Unleached synthetic 2-line ferrihydrite XRD spectrum. CuK $\alpha$  radiation with monochromator, Cardiff University, UK.*



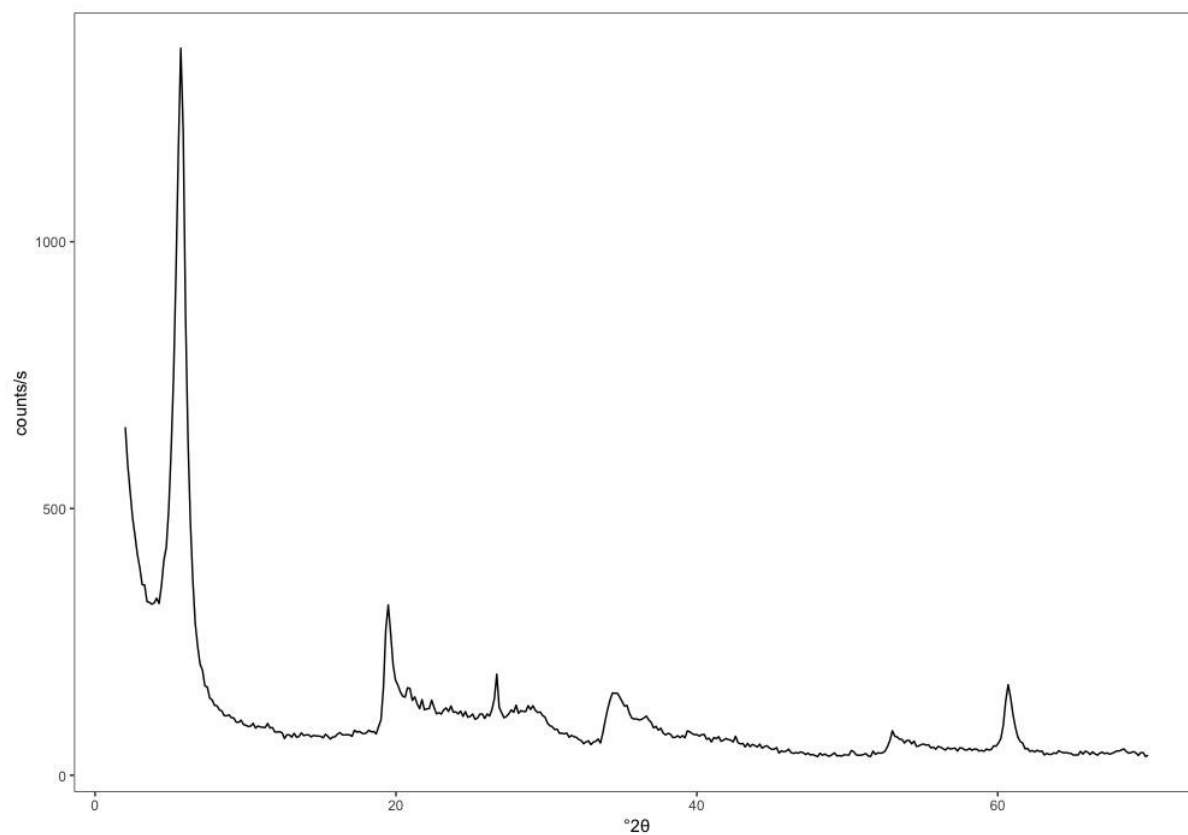
*Figure A3: Unleached synthetic goethite XRD spectrum. CuK $\alpha$  radiation without monochromator, National Museums Collection Centre, Edinburgh, UK.*



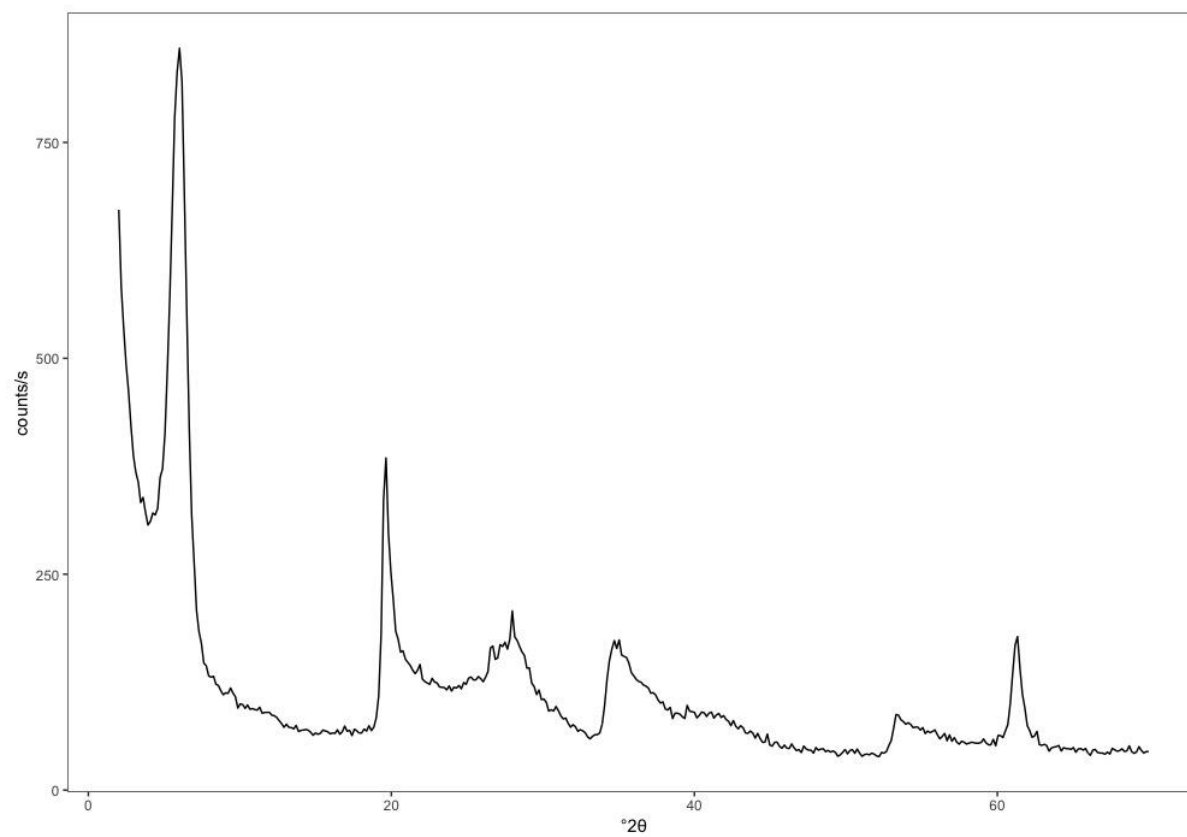
*Figure A4: Unleached natural UoE goethite XRD spectrum. CuK $\alpha$  radiation without monochromator, University of Edinburgh, UK.*



*Figure A5: Unleached synthetic magnetite XRD spectrum. CuK $\alpha$  radiation without monochromator, National Museums Collection Centre, Edinburgh, UK.*



*Figure A6: Unleached natural NMS nontronite XRD spectrum. CuK $\alpha$  radiation with monochromator, Cardiff University, UK.*



*Figure A7: Unleached natural NAu-2 nontronite XRD spectrum. CuK $\alpha$  radiation with monochromator, Cardiff University, UK.*



## Appendix B to

**The use of operationally-defined sequential Fe extraction methods for mineralogical applications: a cautionary tale from Mössbauer spectroscopy**

*Laura E. Hepburn, Ian B. Butler, Adrian Boyce, Christian Schröder*

**Content**

**Figure B8:** Observed and fitted Mössbauer spectroscopy spectrum for the unleached natural Ivigtut siderite sample used in this study.

**Figure B9:** Observed and fitted Mössbauer spectroscopy spectrum for the unleached synthetic 2-line ferrihydrite sample used in this study.

**Figure B10:** Observed and fitted Mössbauer spectroscopy spectrum for the unleached synthetic goethite sample used in this study.

**Figure B11:** Observed and fitted Mössbauer spectroscopy spectrum for the unleached natural UoE goethite sample used in this study.

**Figure B12:** Observed and fitted Mössbauer spectroscopy spectrum for the unleached synthetic magnetite sample used in this study.

**Figure B13:** Observed and fitted Mössbauer spectroscopy spectrum for the unleached natural NMS nontronite sample used in this study.

**Figure B14:** Observed and fitted Mössbauer spectroscopy spectrum for the unleached natural NAu-2 nontronite sample used in this study.

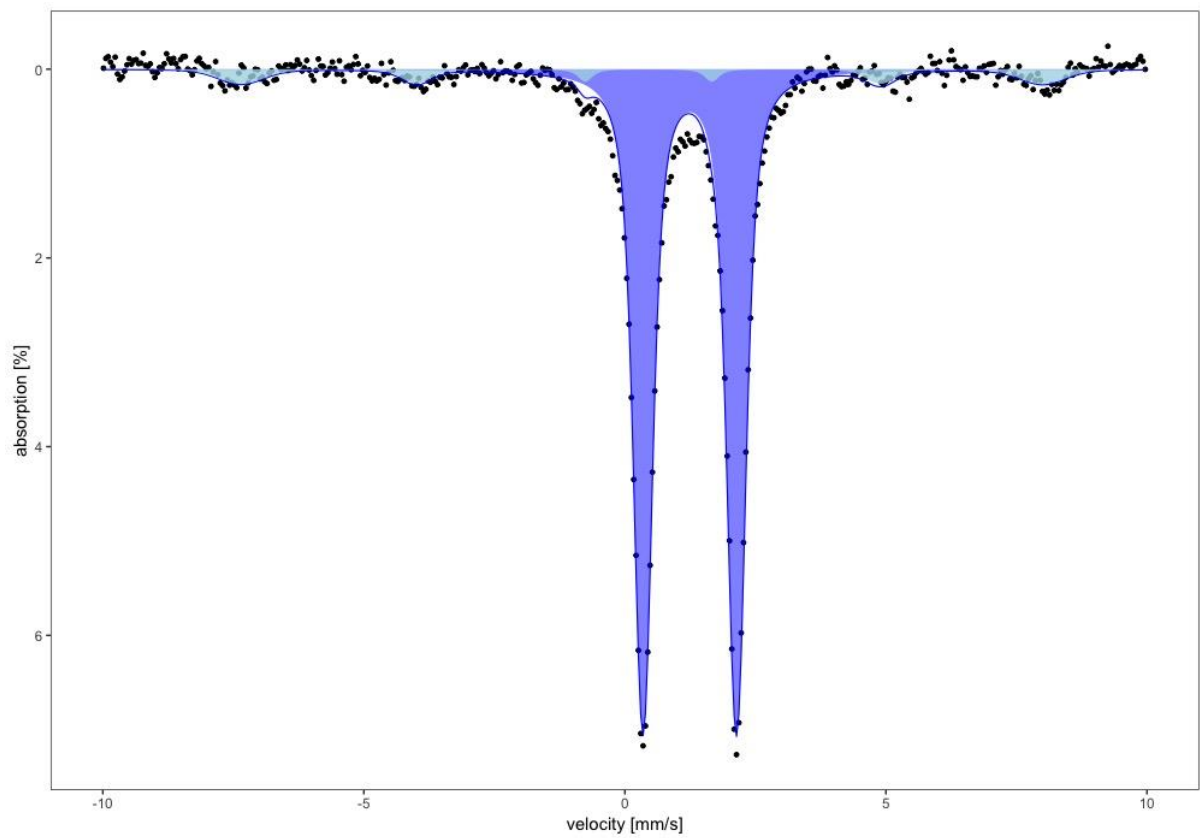


Figure B8: Observed (black circles) and fitted (blue line) Mössbauer spectroscopy spectrum for the unleached natural Ivigtut siderite sample used in this study. The dark blue-filled area is the typical  $\text{Fe}^{2+}$  siderite signature. The light blue-filled area represents  $\text{Fe}^{3+}$  in an iron oxide phase of low crystallinity and shows that a small amount of the iron in Ivigtut siderite is oxidised.

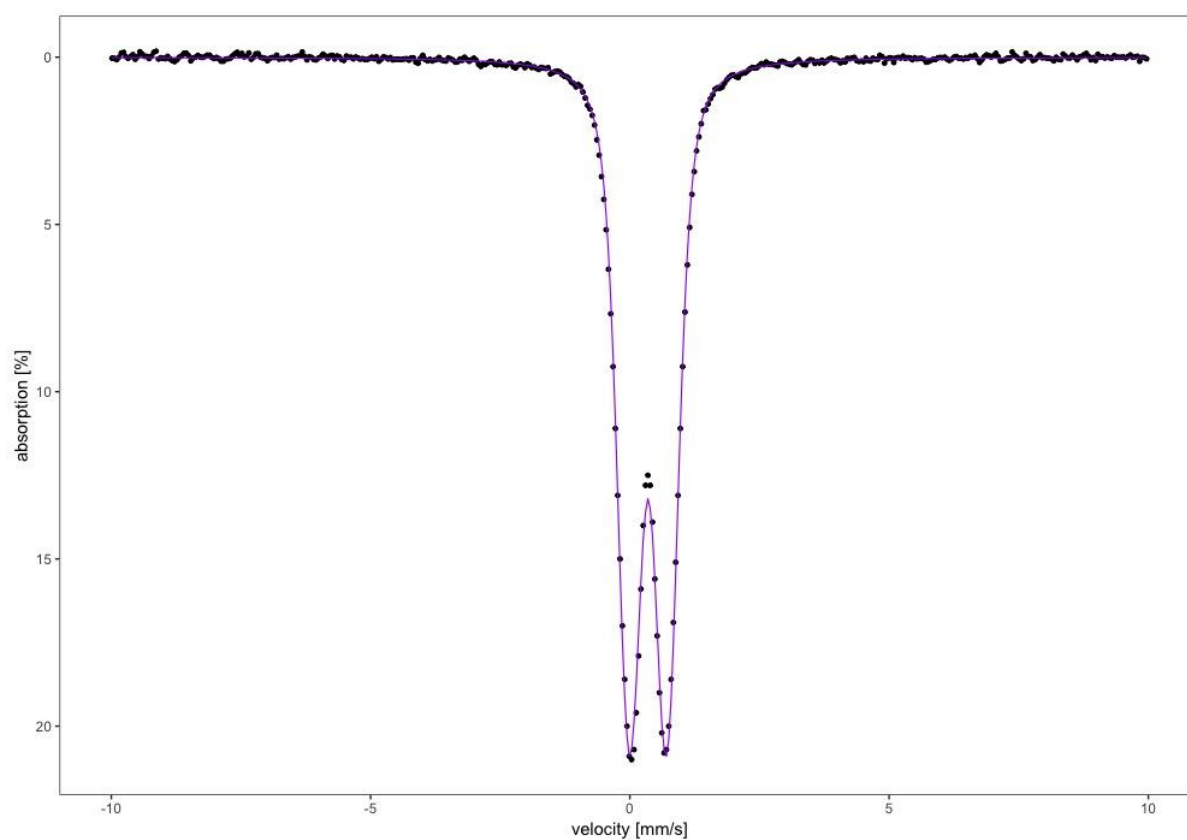


Figure B9: Observed (black circles) and fitted (purple line) Mössbauer spectroscopy spectrum for the unleached synthetic 2-line ferrihydrite sample used in this study.

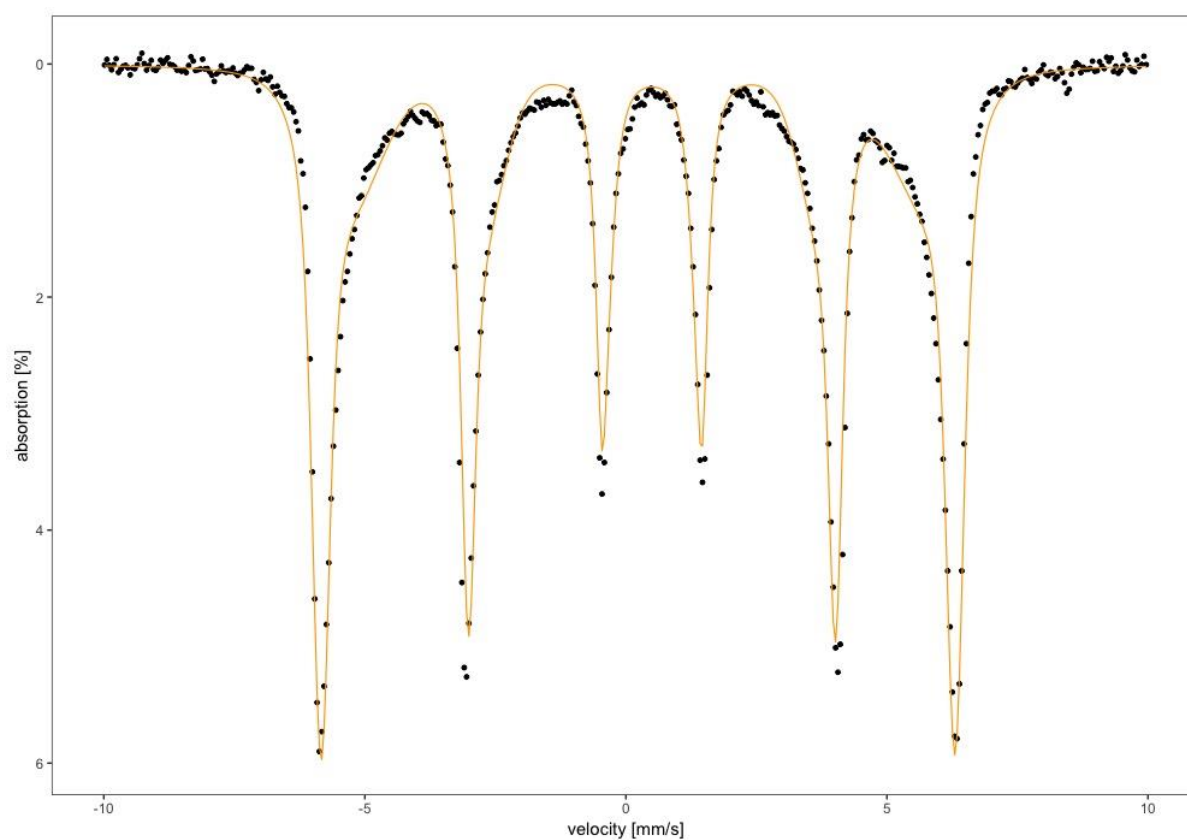


Figure B10: Observed (black circles) and fitted (orange line) Mössbauer spectroscopy spectrum for the unleached synthetic goethite sample used in this study.

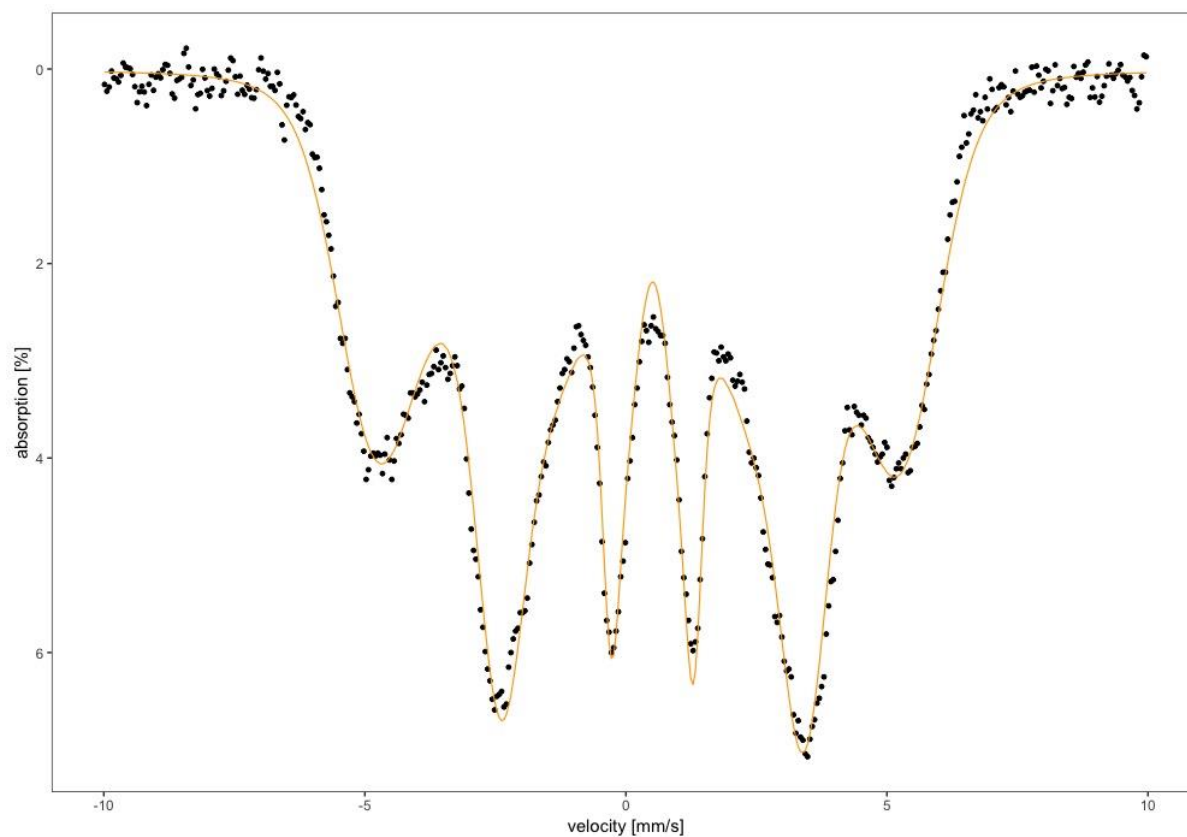


Figure B11: Observed (black circles) and fitted (orange line) Mössbauer spectroscopy spectrum for the unleached natural UoE goethite sample used in this study.

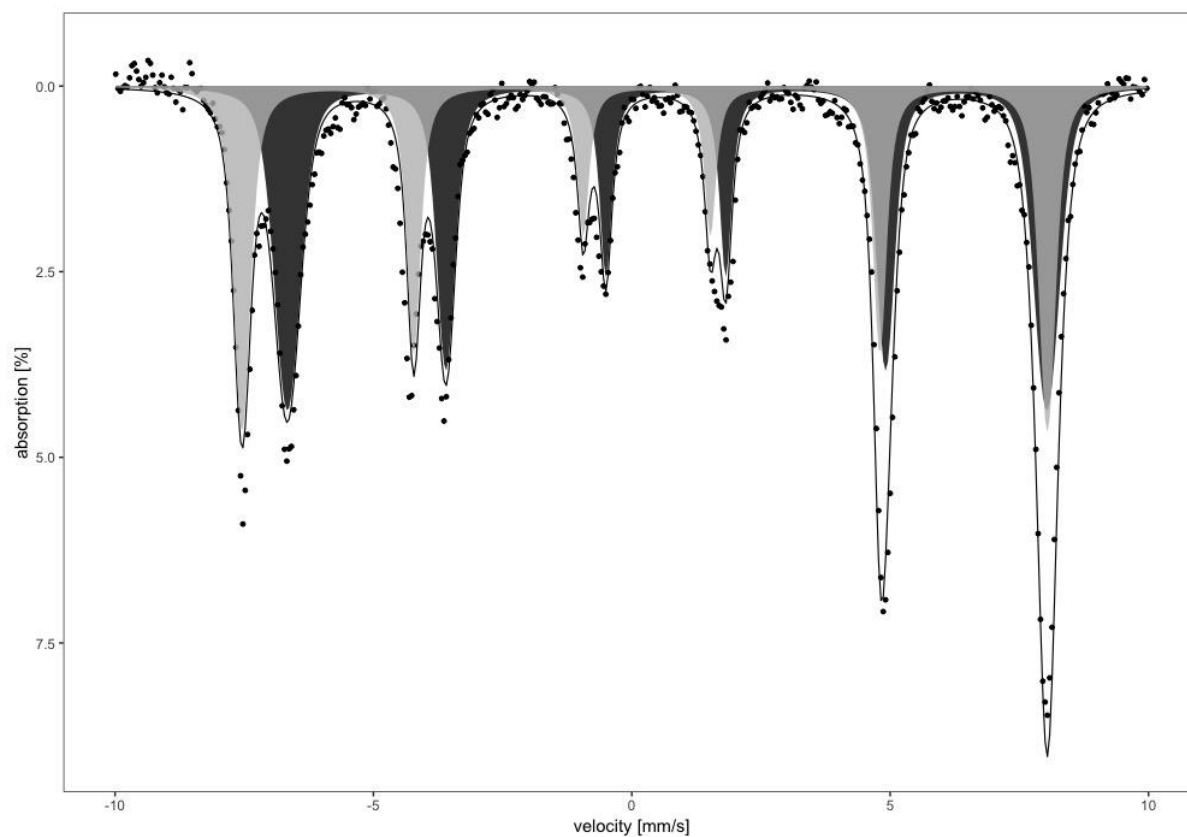


Figure B12: Observed (black circles) and fitted (black line) Mössbauer spectroscopy spectrum for the unleached synthetic magnetite sample used in this study. Magnetite is represented by two subspectra. The black-filled area represents  $\text{Fe}^{3+}$  in tetrahedral crystal lattice sites. The grey-filled area represents  $\text{Fe}^{2+}$  and  $\text{Fe}^{3+}$  in octahedral lattice sites with an average oxidation state of  $\text{Fe}^{2.5+}$ .

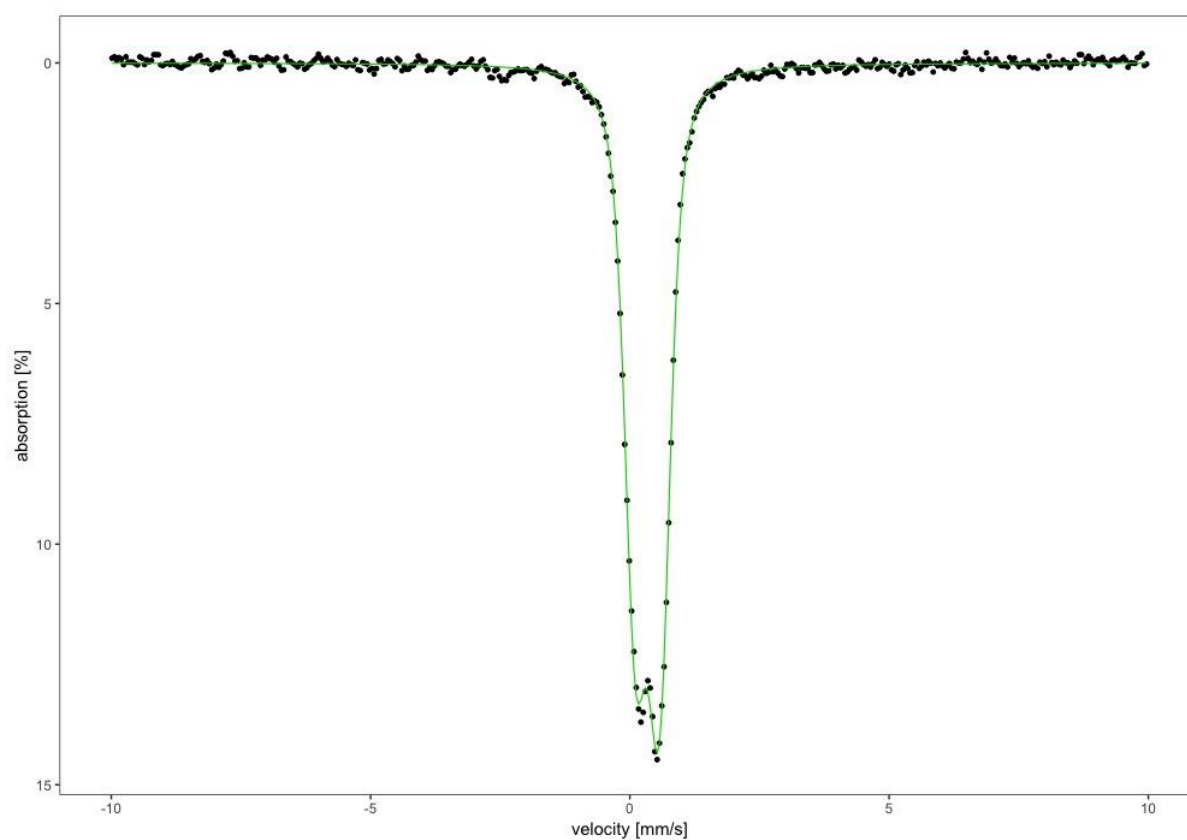


Figure B13: Observed (black circles) and fitted (green line) Mössbauer spectroscopy spectrum for the unleached natural NMS nontronite sample used in this study.

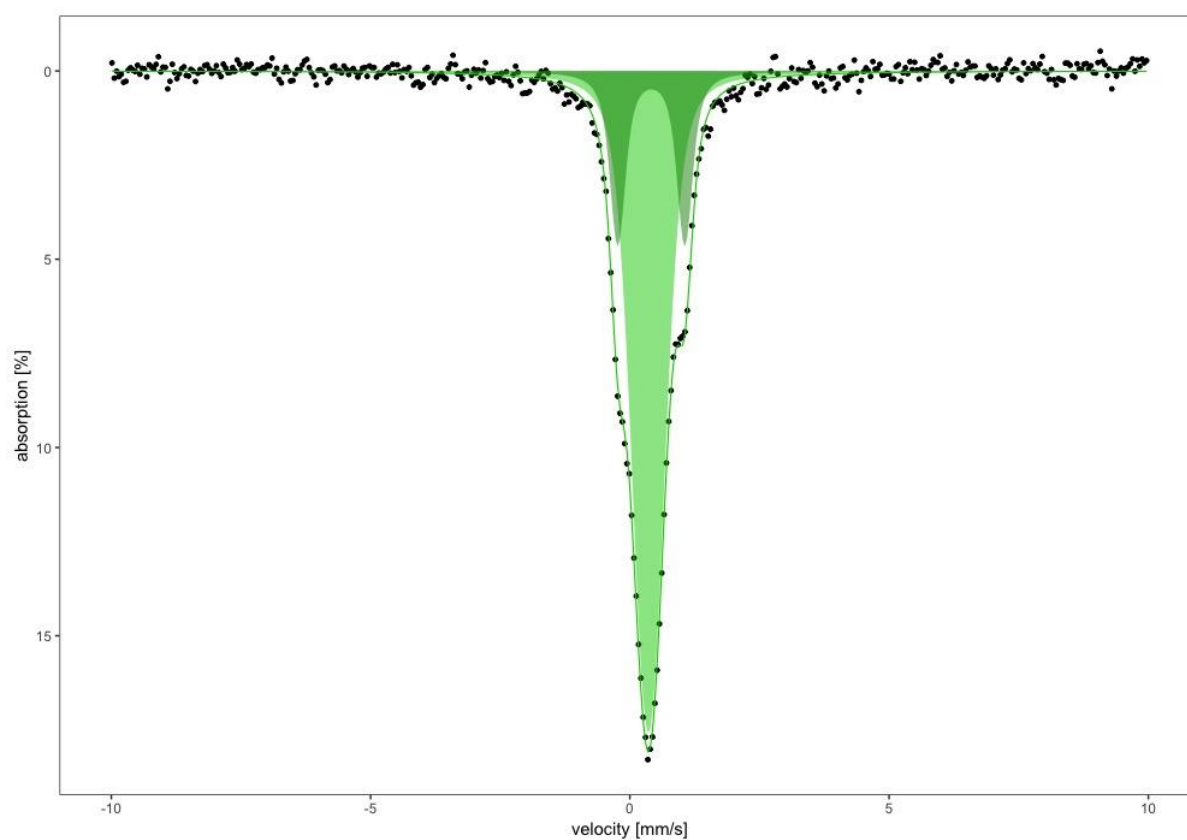


Figure B14: Observed (black circles) and fitted (green line) Mössbauer spectroscopy spectrum for the unleached natural NAu-2 nontronite sample used in this study. The Mössbauer parameters distinguish Fe<sup>3+</sup> in a tetrahedral position in the crystal lattice (light green-filled area) from Fe<sup>3+</sup> in an octahedral position (dark green-filled area).



## Appendix C to

**The use of operationally-defined sequential Fe extraction methods for mineralogical applications: a cautionary tale from Mössbauer spectroscopy**

*Laura E. Hepburn, Ian B. Butler, Adrian Boyce, Christian Schröder*

**Content**

**Table C1:** Mössbauer spectroscopy parameters for the individual Fe minerals unleached and following their target extraction stage only.

**Table C2:** Mössbauer spectroscopy parameters for the individual Fe minerals unleached and at each stage of the sequential extraction until complete removal was observed.

**Table C3:** Mössbauer spectroscopy parameters for the mixed Fe mineral sample CARB-2, unleached and after each stage of the sequential extraction between  $\text{Fe}_{\text{aca-48}}$  and  $\text{Fe}_{\text{oxa}}$ .

**Table C4:** Mössbauer spectroscopy parameters for the mixed Fe mineral sample CARB-3, unleached and after each stage of the sequential extraction between  $\text{Fe}_{\text{aca-48}}$  and  $\text{Fe}_{\text{HCl-1min}}$ .

**Table C1:** Mössbauer spectroscopy parameters for the individual Fe minerals unleached and following their target extraction stage only. No data is shown for 2-line ferrihydrite since it was completely removed during its target,  $Fe_{hyom}$  extraction. Parameters shown are Recoil compiled site properties.  $\delta$  = isomer shift;  $\Delta E_Q$  = quadrupole splitting;  $B_{hf}$  = internal magnetic field;  $n$  = natural;  $s$  = synthetic.

Fe mineral	Ext. stage	Site	$\delta$ <i>mm/s</i>	$\Delta E_Q$ <i>mm/s</i>	$B_{hf}$ <i>T</i>
Ivigtut siderite ( <i>n</i> )	$Fe_{UL}$	$Fe^{2+}$	1.24	1.79	
		$Fe^{3+}$	0.39	-0.06	47.6
	$Fe_{aca-48}$	$Fe^{2+}$	1.24	1.78	
goethite ( <i>s</i> )	$Fe_{UL}$	$Fe^{3+}$	0.33	-0.17	37.7
	$Fe_{di-ct}$	$Fe^{3+}$	0.51	-0.22	37.8
UoE goethite ( <i>n</i> )	$Fe_{UL}$	$Fe^{3+}$	0.41	-0.12	31.3
	$Fe_{di-ct}$	$Fe^{3+}$	0.44	-0.07	32.3
magnetite ( <i>s</i> )	$Fe_{UL}$	$Fe^{2.5+}$	0.67	0.01	45.6
		$Fe^{3+}$	0.27	-0.02	48.3
	$Fe_{oxa}$	$Fe^{2.5+}$	0.69	0.01	45.9
		$Fe^{3+}$	0.26	-0.03	48.4
NMS nontronite ( <i>n</i> )	$Fe_{UL}$	$Fe^{3+}$	0.34	0.46	
	$Fe_{HCl-1min}$	$Fe^{3+}$	0.37	0.36	
NAu-2 nontronite ( <i>n</i> )	$Fe_{UL}$	$Fe^{3+}$ ( <i>tetra</i> )	0.36	0.10	
		$Fe^{3+}$ ( <i>octa</i> )	0.41	1.3	
	$Fe_{HCl-1min}$	$Fe^{3+}$ ( <i>tetra</i> )	0.37	0.00	

**Table C2:** Mössbauer spectroscopy parameters for the individual Fe minerals unleached and at each stage of the sequential extraction until complete removal was observed. Parameters shown are Recoil compiled site properties.  $\delta$  = isomer shift;  $\Delta E_Q$  = quadrupole splitting;  $B_{hf}$  = internal magnetic field; n = natural; s = synthetic.

Fe mineral	Ext. stage	Site	$\delta$ mm/s	$\Delta E_Q$ mm/s	$B_{hf}$ T
Ivigtut siderite (n)	Fe <sub>UL</sub>	Fe <sup>2+</sup>	1.24	1.79	47.6
		Fe <sup>3+</sup>	0.39	-0.06	
	Fe <sub>aca-48</sub>	Fe <sup>2+</sup>	1.24	1.78	
	Fe <sub>hyam</sub>	Fe <sup>2+</sup>	1.25	1.80	
	Fe <sub>di-ct</sub>	Fe <sup>2+</sup>	1.25	1.79	
	Fe <sub>oxa</sub>	Fe <sup>2+</sup>	1.25	1.79	
2-line ferrihydrite (s)	Fe <sub>UL</sub>	Fe <sup>2+</sup>	0.35	0.72	
	Fe <sub>aca-48</sub>	Fe <sup>2+</sup>	0.35	0.77	
goethite (s)	Fe <sub>UL</sub>	Fe <sup>3+</sup>	0.33	-0.17	37.7
	Fe <sub>aca-48</sub>	Fe <sup>3+</sup>	0.44	-0.08	37.6
	Fe <sub>hyam</sub>	Fe <sup>3+</sup>	0.39	-0.16	37.7
UoE goethite (n)	Fe <sub>UL</sub>	Fe <sup>3+</sup>	0.41	-0.12	31.3
	Fe <sub>aca-48</sub>	Fe <sup>3+</sup>	0.37	-0.12	31.8
	Fe <sub>hyam</sub>	Fe <sup>3+</sup>	0.34	-0.16	32.0
	Fe <sub>di-ct</sub>	Fe <sup>3+</sup>	0.38	-0.12	31.5
	Fe <sub>oxa</sub>	Fe <sup>3+</sup>	0.33	-0.17	31.3
	Fe <sub>HCl-1min</sub>	Fe <sup>3+</sup>	0.34	-0.16	31.0
magnetite (s)	Fe <sub>UL</sub>	Fe <sup>2.5+</sup>	0.67	0.01	45.6
		Fe <sup>3+</sup>	0.27	-0.02	48.3
	Fe <sub>aca-48</sub>	Fe <sup>2.5+</sup>	0.69	0.02	46.1
		Fe <sup>3+</sup>	0.25	-0.03	48.4

	$\text{Fe}_{\text{hyam}}$	$\text{Fe}^{2.5+}$	0.69	0.02	46.1
		$\text{Fe}^{3+}$	0.26	-0.03	48.7
	$\text{Fe}_{\text{di-ct}}$	$\text{Fe}^{2.5+}$	0.69	0.02	46.0
		$\text{Fe}^{3+}$	0.27	-0.03	48.6
NMS nontronite ( <i>n</i> )	$\text{Fe}_{\text{UL}}$	$\text{Fe}^{3+}$	0.34	0.46	
	$\text{Fe}_{\text{aca-48}}$	$\text{Fe}^{3+}$	0.33	0.48	
	$\text{Fe}_{\text{hyam}}$	$\text{Fe}^{3+}$	0.33	0.46	
	$\text{Fe}_{\text{di-ct}}$	$\text{Fe}^{3+}$	0.37	0.67	
	$\text{Fe}_{\text{oxa}}$	$\text{Fe}^{3+}$	0.21	0.67	
	$\text{Fe}_{\text{HCl-1min}}$	$\text{Fe}^{3+}$	0.28	0.00	
NAu-2 nontronite ( <i>n</i> )	$\text{Fe}_{\text{UL}}$	$\text{Fe}^{3+} (\text{tetra})$	0.36	0.10	
		$\text{Fe}^{3+} (\text{octa})$	0.41	1.3	
	$\text{Fe}_{\text{aca-48}}$	$\text{Fe}^{3+} (\text{tetra})$	0.38	0.00	
	$\text{Fe}_{\text{hyam}}$	$\text{Fe}^{3+} (\text{tetra})$	0.37	0.00	
	$\text{Fe}_{\text{di-ct}}$	$\text{Fe}^{3+} (\text{tetra})$	0.36	0.00	
	$\text{Fe}_{\text{oxa}}$	$\text{Fe}^{3+} (\text{tetra})$	0.38	0.00	
	$\text{Fe}_{\text{HCl-1min}}$	$\text{Fe}^{3+} (\text{tetra})$	0.36	0.00	

**Table C3:** Mössbauer spectroscopy parameters for the mixed Fe mineral sample CARB-2, unleached and after each stage of the sequential extraction between  $Fe_{aca-48}$  and  $Fe_{oxa}$ ; all trace of Fe was removed during the  $Fe_{HCl-1min}$  extraction. Parameters shown are Recoil compiled site properties.  $\delta$  = isomer shift;  $\Delta E_Q$  = quadrupole splitting;  $B_{hf}$  = internal magnetic field; n = natural; s = synthetic.

Ext. stage	Fe mineral	Site	$\delta$ mm/s	$\Delta E_Q$ mm/s	$B_{hf}$ T
CARB-2 $Fe_{UL}$	Ivigtut siderite (n)	$Fe^{2+}$	1.24	1.79	
	2-line ferrihydrite (s)	$Fe^{2+}$	0.35	0.72	
	goethite (s)	$Fe^{3+}$	0.33	-0.17	37.7
	magnetite (s)	$Fe^{2.5+}$	0.67	0.01	45.6
		$Fe^{3+}$	0.27	-0.02	48.3
	NMS nontronite (n)	$Fe^{3+}$	0.34	0.46	
CARB-2 $Fe_{aca-48}$	Ivigtut siderite (n)	$Fe^{2+}$	1.24	1.79	
	2-line ferrihydrite (s)	$Fe^{2+}$	0.35	0.72	
	goethite (s)	$Fe^{3+}$	0.33	-0.17	37.7
	magnetite (s)	$Fe^{2.5+}$	0.67	0.01	45.6
		$Fe^{3+}$	0.27	-0.02	48.3
	NMS nontronite (n)	$Fe^{3+}$	0.34	0.46	
CARB-2 $Fe_{hyam}$	Ivigtut siderite (n)	$Fe^{2+}$	1.24	1.79	
	goethite (s)	$Fe^{3+}$	0.33	-0.17	37.7
	magnetite (s)	$Fe^{2.5+}$	0.67	0.01	45.6
		$Fe^{3+}$	0.27	-0.02	48.3
	NMS nontronite (n)	$Fe^{3+}$	0.34	0.46	
CARB-2 $Fe_{di-ct}$	Ivigtut siderite (n)	$Fe^{2+}$	1.24	1.79	
	goethite (s)	$Fe^{3+}$	0.33	-0.17	37.7
	magnetite (s)	$Fe^{2.5+}$	0.67	0.01	45.6

		Fe <sup>3+</sup>	0.27	-0.02	48.3
	NMS nontronite ( <i>n</i> )	Fe <sup>3+</sup>	0.34	0.46	
CARB-2 Fe <sub>oxa</sub>	Ivigtut siderite ( <i>n</i> )	Fe <sup>2+</sup>	1.24	1.79	
	goethite ( <i>s</i> )	Fe <sup>3+</sup>	0.33	-0.17	37.7

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**Table C4:** Mössbauer spectroscopy parameters for the mixed Fe mineral sample CARB-3, unleached and after each stage of the sequential extraction between  $Fe_{aca-48}$  and  $Fe_{HCl-1min}$ ; all trace of Fe was removed during the final  $Fe_U$  extraction. Parameters shown are Recoil compiled site properties.  $\delta$  = isomer shift;  $\Delta E_Q$  = quadrupole splitting;  $B_{hf}$  = internal magnetic field;  $n$  = natural;  $s$  = synthetic.

Ext. stage	Fe mineral	Site	$\delta$ mm/s	$\Delta E_Q$ mm/s	$B_{hf}$ T
CARB-3 $Fe_U$	Ivigtut siderite ( <i>n</i> )	$Fe^{2+}$	1.24	1.79	
	2-line ferrihydrite ( <i>s</i> )	$Fe^{2+}$	0.35	0.72	
	UoE goethite ( <i>n</i> )	$Fe^{3+}$	0.41	-0.12	31.3
	magnetite ( <i>s</i> )	$Fe^{2.5+}$	0.67	0.01	45.6
		$Fe^{3+}$	0.27	-0.02	48.3
	NAu-2 nontronite ( <i>n</i> )	$Fe^{3+}$ ( <i>tetra</i> )	0.36	0.12	
		$Fe^{3+}$ ( <i>octa</i> )	0.41	0.16	
CARB-3 $Fe_{aca-48}$	Ivigtut siderite ( <i>n</i> )	$Fe^{2+}$	1.24	1.79	
	2-line ferrihydrite ( <i>s</i> )	$Fe^{2+}$	0.35	0.72	
	UoE goethite ( <i>n</i> )	$Fe^{3+}$	0.41	-0.12	31.3
	magnetite ( <i>s</i> )	$Fe^{2.5+}$	0.67	0.01	45.6
		$Fe^{3+}$	0.27	-0.02	48.3
	NAu-2 nontronite ( <i>n</i> )	$Fe^{3+}$ ( <i>tetra</i> )	0.36	0.12	
		$Fe^{3+}$ ( <i>octa</i> )	0.41	0.16	
CARB-3 $Fe_{hyam}$	Ivigtut siderite ( <i>n</i> )	$Fe^{2+}$	1.24	1.79	
	UoE goethite ( <i>n</i> )	$Fe^{3+}$	0.41	-0.12	31.3
	magnetite ( <i>s</i> )	$Fe^{2.5+}$	0.67	0.01	45.6
		$Fe^{3+}$	0.27	-0.02	48.3
	NAu-2 nontronite ( <i>n</i> )	$Fe^{3+}$ ( <i>tetra</i> )	0.36	0.12	
		$Fe^{3+}$ ( <i>octa</i> )	0.41	1.29	

CARB-3 Fe <sub>di-ct</sub>	Ivigtut siderite ( <i>n</i> )	Fe <sup>2+</sup>	1.24	1.79	
	UoE goethite ( <i>n</i> )	Fe <sup>3+</sup>	0.41	-0.12	31.3
	magnetite ( <i>s</i> )	Fe <sup>2.5+</sup>	0.67	0.01	45.6
		Fe <sup>3+</sup>	0.27	-0.02	48.3
	NAu-2 nontronite ( <i>n</i> )	Fe <sup>3+</sup> ( <i>tetra</i> )	0.36	0.12	
CARB-3 Fe <sub>oxa</sub>	Ivigtut siderite ( <i>n</i> )	Fe <sup>2+</sup>	1.24	1.79	
	UoE goethite ( <i>n</i> )	Fe <sup>3+</sup>	0.41	-0.12	31.3
	NAu-2 nontronite ( <i>n</i> )	Fe <sup>3+</sup> ( <i>tetra</i> )	0.36	0.12	
CARB-3 Fe <sub>HCl-1min</sub>	Ivigtut siderite ( <i>n</i> )	Fe <sup>2+</sup>	1.24	1.79	
	NAu-2 nontronite ( <i>n</i> )	Fe <sup>3+</sup> ( <i>tetra</i> )	0.36	0.12	

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